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VPATEWGTSQASGDGGPYFDEADVWIEFLNENNISWANWSLTNKNEVSGAFTPFELGKSN
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     The invention relates to a mutant alkaline cellulase derived from the Bacillus sp. KSM-S237 akaline cellulase Egl-237 (ADE71407). The mutant enzyme is created by delating one or more amino acid residues between residues 343-373 of the wild-type enzyme, and then inserting a 2-15 residue peptide into the deletion site. The invention also encompasses a gene encoding a mutant alkaline cellulase of the invention, and vectors and host comprising a mutant alkaline cellulase-encoding gene. The mutant alkaline cellulases of the invention pawe my very close to the ph of laundry water (around ph 10.5) and are therefore useful as enzymes for detergents Sequences ADE71413-ADE71415 represent
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                                                                                                                                                                                                                                                                                                                                                                                                                       from the 343-
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                                                                                                                                                                                                                                                                                                                                                                                                    cellulase for use as an enzyme for detergents is thing one or more amino acid residue groups from the 34 th SEQ ID No:1 and then inserting a peptide into the
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   167 PHIIYELANEPSSNNNGGAGIPNNEEGWKAVKEYADPIVQMLKKSGNADDN
                                                                                                                                       Alkaline cellulase; Egl-N131b; detergent; laundry; enzyme
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                                                                                                            (llus sp. KSM-N131 alkaline cellulase Egl-N131b
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       alkaline cellulases from other Badillus species.
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                        ADE71415 standard; protein; 810 AA.
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to 373-positions in
deletion site.
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287 VPATEWGTSQANGDGGPYPDEADVWIEFLNENNISWANWSLINKNEVSGAFTPFELGKSN
                                                              361 ATNLDPGPDHVWAPEBLSLSGBYVRARIKGVNYEPIDRTKYTKVLWDFNDGTKQGFGVNS
                                                                                                                               ELIAVDNENNTLKVSGLDVSNDVSDGNFWANARLSANGWGKSVDILGAEKLTMDV
                                                                                                                                                                         VAIAAIPQSSKSGWANPERAVRVNAEDFVQQTDGKYKAGLTITGEDAPNLKNI
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                                                                                                         SPNKELIAVDNENNTLKVSGLDVSNDVSDGNFWANARLSANGWGKSVDILGAEKLTMDV
                                                                                                                                                                                                                                        LEVGTDAADVIYLDNIKVIGTEVBIPVVHDPKGRAVLPSVFBDGTRQ
                                                                                                                                                                                                                                                                                                                                    587 GWDWAGESGVKTALTIEEANOGNALSWEFGYPEVKPSDNWATAPRLDFWKSDLVRGENDY
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Location/Qualifiers
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAG80267 standard; protein; 813 AA.
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N-PSDB; AAI69288.
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Enzyme, endo-beta-1,4-glucanase, detergent, textile finishing process, oil inquetry, biomass degradation, laundry, stone washing; EC 3.2.1.4; pulp processing, animal feed.

AA349 strain DSM 12648. Location/Qualifiers

Bacillus

Barillus sp. endo-beta-1,4-glucanase.

(first entry)

(revised)

23-OCT-2003 07-MAY-2003 ABG76403;

340. .540 | Jabel = Cellulase binding site | note= "This site is claimed in claim 24"

WO200299091-A2;

12-DEC-2002.

note= "Encoded by GAR"

Misc-differend Binding-site

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sp. The alkaline cellulase gene is used for the preparation of an alkaline cellulase useful as a textile detergent and a textile treating agent. This sequence represents the Bacillus sp. alkaline cellulase N131b described in the method of the invention
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  388 IKGVNYEPIDRIKYTKVLMDFNDGTKQGFGVNSDSPNKELIAVDNENNTLKVSGLDVSND 447
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   437 VSDGNFWANARLSANGWGKSVDILGAEKLIMDVIVDEPITVAIAAIPQSSKSGWANPERA 496
                                                                                                                                                                                                                                      on describes a novel alkaline cellulase gene from a Bacillus
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    317 FLNENNISWANWSLTINGNEVSGAFTPFELGKSNATSLDPGPDQVWYPEELSLSGETVRAR
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                                                                                                                                                                                                                                                                                       WPPEILNDNAYKALSNDWDSNMIRLAMYVGENGYATNPELIKQRVIDGIELALENDMYVI
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                                                                                                                                                                                      Gaps
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                                                                                                                                                 94.6%; Score 4123; DB 5; Length 813; 97.1%; Pred. No. 6e-273; ive 10; Mismatches 13; Indels
                                                                                                                                                                 Best Local Similarity 97.18
Matches 774; Conservative
                                                                                                                   Sequence 813 AA;
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activity (EC 3.2.1.4), comprising: (a) a polypeptide encoded by the DNA sequence appearing as ABX11841; (b) a polypeptide produced by culturing cell comprising man and sequence under conditions where the DNA sequence is expressed; (c) an endo-beta-1, 4-glucanase enzyme having at least 978 sequence identity to the amino acid sequence appearing as ABG76403; or (d) a polypeptide having endo-beta-1, 4-glucanase activity that is encode by a polymucleotide that hybridizes to the DNA under hybridisation conditions comprising 5X SC at 45 plusoc and washing conditions comprising 2X SC at 60 plusoc. Also included are the encoding DNA ements improved performance in pulp processing, textile treatment, laundry processes, extraction processes or in animal feed. The present sequence represents the endo-beta-1,4-glucanase. (Updated on 23-OCT-2003 to standardise OS field) sequence finishing processes, oil industry, blomass degradation, laundry and st washing. The invention provides enzymes having substantial beta-1,4-glucanase activity under slightly acid to alkaline conditions and an expression vector (comprising the DNA gequen an expression vector (comprising the following operably linked alemen a transcription promoter, a DNA segment encoding the enzyme and transcription terminator), a cultured cell comprising the vector and expressing the enzyme, a method for degradation of cellulose-contain biomass that is treated with the enzyme or enzyme composition cited and and a hybrid endo-glucanase (exhibiting endo- beta-1,4-glucanase acti comprising the cellulase binding domain, CBD, of the enzyme, and a catalytic domain (CAD) from sources other than Bacillus sp. A349 str DSM12648). The enzymes are useful in detergent composition, textile The invention relates to an enzyme exhibiting endo-Claim 1; Page 45-48; 51pp; English

949

617 EPGYPEVKPSDNWATAPRLDFWKSDLVRGENDYVTPDFYLDPVRATEGAMNINLVPQPPT

628 BFGYPEVKPSDNWATAPRLDFWKSDLVRGENDYVAPDFYLDPVRATEGAMNINLVFQPPT

ð 셤 ò 677 NGYWYQAPKTYTINPDELERANQVNGLYHYEVKINVRDITNIQDDTLLRNWMIIFADVES

NGYWVQAPKTYTINFDELEEANQVNGLYHYEVKINVRDITNIQDDTLLRNMMI1PADVES

688

747

807

737 DFAGRVFVDNYRFBGAATTBFVEPEPVDFGEETPFVDEKBAKKEQKEAEKEEKKEKKER 796

748 DFAGRVEVDNVRFEGAATTEPVEPEPVDPGEETPPVDEKEAKKEQKEAEKEEKEAVKEEK

ò 셤

activity, useful in finishing processes, biomass

New enzyme exhibiting endo-beta-1,4-glucanabe detergent compositions, oil industry textile degradation, laundry, and stone washing.

Gibson K;

Eskelund

Outtrup H, Schuelein M,

WPI; 2003-256232/25. N-PSDB; ABX11841

(NOVO) NOVOZYMES AS

06-JUN-2001; 2001DK-00000879 06-JUN-2002; 2002WO-DK000381

Sequence 773 AA,

93.1%; Score 4059;

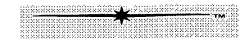
Query Match

46403 standard; protein; 773 AA

808 KBAKEEKKAVKNEAKKK 824

797

6; Length 773;



MACHINE-ASSISTED TRANSLATION (MAT):

(19)【発行国】 (19)[ISSUING COUNTRY] 日本国特許庁(JP) Japan Patent Office (JP)

(12)【公報種別】 (12)[GAZETTE CATEGORY] 公開特許公報(A) Laid-open Kokai Patent (A)

(11)【公開番号】 (11)[KOKAI NUMBER]

2001-231569(P2001-231569A) 2001-231569(P2001-231569A)

(43)【公開日】(43)[DATE OF FIRST PUBLICATION]平成13年8月28日(200 August 28, Heisei 13 (2001. 8.28)

1. 8. 28)

(54)【発明の名称】 (54)[TITLE OF THE INVENTION]

アルカリセルラーゼ遺伝子 Alkali cellulase gene

(51)【国際特許分類第 7 版】 (51)[IPC 7]
C12N 15/09 ZNA C12N 15/09 ZNA
1/00 1/15 1/15

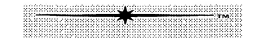
1/191/191/211/215/105/109/429/42

[FI] [FI] C12N 1/00 T C12N 1/00

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 1/19

Т



1/21 1/21

9/42 9/42

15/00 ZNA A 15/00 ZNA A 5/00 A

【審査請求】 未請求 [REQUEST FOR EXAMINATION] No

【請求項の数】 6 [NUMBER OF CLAIMS] 6

【出願形態】 O L [FORM OF APPLICATION] Electronic

【全頁数】 22 [NUMBER OF PAGES] 22

(21)【出願番号】 (21)[APPLICATION NUMBER]

2000-47237(P2000-47237) 2000-47237(P2000-47237)

(22)【出願日】 (22)[DATE OF FILING]

平成12年2月24日 (200 February 24, Heisei 12 (2000. 2.24)

0.2.24

(71)【出願人】 (71)[PATENTEE/ASSIGNEE]

【識別番号】[ID CODE]000000918000000918

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目14番10号

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100068700

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【テーマコード(参考)】

[THEME CODE (REFERENCE)]

4B024

4B024

4B050

4B050

4B065

4B065

【Fターム(参考)】

[F TERM (REFERENCE)]

4B024 BA11 CA04 CA09 DA07 4B024 BA11 CA04 CA09 DA07 EA04 GA11

EA04 GA11 GA19 GA27 HA01 GA19 GA27 HA01 HA19

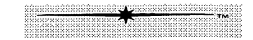
HA19

4B050 CC03 DD02 LL04

4B050 CC03 DD02 LL04

4B065 AA15X AA15Y AB01 BA02 BA22 CA31

4B065 AA15X AA15Y AB01 CA57



BA02 BA22 CA31 CA57

(57)【要約】

(修正有)

【解決手段】

特定の配列を有する2種類の アミノ酸配列のいずれか、又は 個のアミノ酸が欠失、置換若し くは付加されたアミノ酸配列を コードするアルカリセルラーゼ 遺伝子、組換えベクター及び形 質転換体。

【効果】

剤、繊維処理剤等として有用な アルカリセルラーゼを単一且つ 大量に生産することが可能であ using this gene. る。

【特許請求の範囲】

【請求項1】

配列の1若しくは数個のアミノ 酸が欠失、置換若しくは付加さ れたアミノ酸配列をコードする アルカリセルラーゼ遺伝子。

【請求項2】

塩基配列、又は該塩基配列の1

(57)[ABSTRACT OF THE DISCLOSURE]

(Amendments Included)

[PROBLEM TO BE SOLVED]

The alkali cellulase gene, recombinant vector, and transformed body which code the amino 該アミノ酸配列の1若しくは数 acid sequence of which 1 of either of two kinds of amino acid sequences which has a specific sequence, or this amino acid sequence, or some amino acids were deleted, substituted or added.

[ADVANTAGE]

この遺伝子を用いて衣料用洗 Alkali cellulase useful as the detergent for garments, a fiber processing agent, etc. can be produced individually and in large quantities

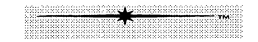
[CLAIMS]

[CLAIM 1]

配列番号1若しくは2に示す The alkali cellulase gene which codes the アミノ酸配列、又は該アミノ酸 amino acid sequence of which the amino acid sequence shown in sequence number 1 or 2, 1 of this amino acid sequence, or some amino acids were deleted, substituted or added.

[CLAIM 2]

配列番号3若しくは4に示す The alkali cellulase gene which has the base acid sequence of which the base sequence 若しくは数個の塩基が欠失、置 shown in sequence number 3 or 4, 1 of this



列を有するアルカリセルラーゼ substituted or added. 遺伝子。

換若しくは付加された塩基酸配 base sequence, or some bases were deleted,

【請求項3】

を含む組換えベクター。

[CLAIM 3]

請求項1又は2記載の遺伝子 The recombinant vector containing the gene of Claim 1 or 2.

【請求項4】

請求項3記載の組換えベクタ The ーを含む形質転換体。

[CLAIM 4]

transformed body containing the recombinant vector of Claim 3.

【請求項5】

記載の形質転換体。

[CLAIM 5]

宿主が微生物である請求項4 The transformed body of Claim 4 whose host is microorganisms.

【請求項6】

するアルカリセルラーゼの製造 法。

[CLAIM 6]

請求項4又は5に記載の形質 A production of the alkali cellulase, which 転換体を培養することを特徴と cultivates the transformed body of Claim 4 or 5.

【発明の詳細な説明】

[DETAILED DESCRIPTION OF THE INVENTION]

[0001]

[0001]

【発明の属する技術分野】

ドする遺伝子に関する。

[TECHNICAL FIELD OF THE INVENTION]

本発明は、洗剤用酵素として有 This invention relates to the gene which codes 用なアルカリセルラーゼをコー alkali cellulase useful as an enzyme for detergents.

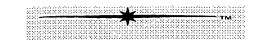
[0002]

[0002]

【従来の技術】

[PRIOR ART]

セルロースは植物細胞壁の主成 A cellulose is the principal component of a



有効利用されるバイオマスの代 表的存在である。セルロースは building material, etc. グルコースが直鎖状に $\beta-1$, 4 結合した巨大分子であるた め、分解によって燃料物質やよ り高付加価値の代謝物質に変換 が可能である。そのためセルロ ースを分解する酵素として、セ ルラーゼ及びその反応産物の有 効利用に関する研究が多岐に行 われている。これらの研究対象 となるセルラーゼは、一般に、 中酸性に最適反応pHを有し、 結晶性セルロースを良好に分解 できる真菌類や嫌気性細菌由来 の酵素が中心となっている。

[0003]

Akiba, Microorganisms, Springer, Berlin, 1982) によって好アルカ リ性バチルス属細菌由来のアル カリセルラーゼが見出されて以 剤への応用が可能となった。そ ルス属細菌の生産するアルカリ セルラーゼ(特公昭60-23 158号公報、特公平6-03 0578号公報、米国特許第4

分で、衣料、紙、建築材料等に plant-cell wall, and is a typical presence of the biomass used effectively for garments, paper, a

> Since the glucose is the macromolecule which carried out the (beta)-1,4 connection linear, the conversion of a cellulose is possible for the fuel matter or a more nearly high-value-added metabolite with a degradation.

> Therefore, as an enzyme which degrades a cellulose, research on an effective usage of cellulase and its reaction production is done variably.

> Generally these the cellulase used as candidates for research has the optimal reaction pH into the in acidity, the enzyme derived from fungi or the anaerophyte which can degrade a crystalline cellulose good has taken the lead.

[0003]

一方、掘越(特公昭 5 0 - 2 8 On the other hand, since the alkali cellulase 515号公報、 Horikoshi & derived from alkali-loving Bacillus bacteria was Alkalophilic discovered by Horikoshi (Japanese Patent Publication No. 50-28515, Horikoshi & Akiba, Alkalophilic Microorganisms, Springer, Berlin, 1982), it has become applicable to the heavy duty detergent for garments of cellulase.

来、セルラーゼの衣料用重質洗 After that, the alkali cellulase (Japanese Patent Publication No. 60-23158, the Japanese Patent の後、実際に好アルカリ性バチ Publication No. 6-030578, US Patent 4945053 grade) which alkali-loving Bacillus bacteria actually produce came to be mixed with the detergent for garments.

The cellulase blending detergent derived from 9 4 5 0 5 3 号等) が衣料用洗 fungi also comes to be marketed after this, it 剤へ配合されるに至った。これ has established the status as an enzyme for



配合洗剤も上市されるようにな lipase, and the amylase. り、プロテアーゼ、リパーゼ、 アミラーゼと並ぶ洗剤用酵素と しての地位を確立してきた。

以降、真菌類由来のセルラーゼ detergents on a par with the protease, the

[0004]

さらに近年、遺伝子工学の発展 るようになっている。アルカリ グ、塩基配列の決定がなされ、 実生産に用いられている例もあ actual production. る。

[0005]

題】

本発明の目的は、洗剤用酵素と 法を確立することにある。

[0006]

【課題を解決するための手段】 カリセルラーゼ生産菌のスクリ

[0004]

Furthermore, production of the enzyme for に伴い、洗剤用酵素の生産も遺 detergents is also mass-produced more gene 伝子組換えにより大量生産され recombinant with development of genetic engineering in recent years.

セルラーゼについても既に数多 As for alkali cellulase, the decision of a cloning くの遺伝子についてクローニン and a base sequence about many genes has already been done, there are examples of

[0005]

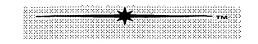
【発明が解決しようとする課 [PROBLEM TO BE SOLVED THE **INVENTION**

There is objective of the invention in して有用なアルカリセルラーゼ establishing the gene which codes alkali をコードする遺伝子及びその遺 cellulase useful as an enzyme for detergents, 伝子を用いた大量かつ単一のア and the method of manufacturing the extensive ルカリセルラーゼを製造する方 and single alkali cellulase using the gene.

[0006]

[MEANS TO SOLVE THE PROBLEM]

本発明者らは、自然界からアル When the present inventors performs a screening of an alkali cellulase producing ーニングを行ったところ、目的 microbe from nature, he discovers the に適う酵素を生産する微生物を microorganisms which produce the enzyme 見出し、さらに当該微生物から which suits the objective, furthermore, by



アルカリセルラーゼをコードする遺伝子をクローン化することにより、本発明を完成した。

[0007]

本発明は、配列番号1若しくは 2に示すアミノ酸配列、又は該 アミノ酸配列の1若しくは数個 のアミノ酸が欠失、置換若しく は付加されたアミノ酸配列をコ ードするアルカリセルラーゼ遺 伝子を提供するものである。ま た、本発明は、配列番号3若し くは4に示す塩基配列、又は該 塩基配列の1若しくは数個の塩 基が欠失、置換若しくは付加さ れた塩基配列を有するアルカリ セルラーゼ遺伝子を提供するも のである。また、本発明は、上 記のアルカリセルラーゼ遺伝子 を含む組換えベクター、及び該 組換えベクターを含む形質転換 体を提供するものである。また、 本発明は、上記の形質転換体を カリセルラーゼの製造法を提供 するものである。

[0008]

【発明の実施の形態】

本発明の遺伝子は、配列番号1 若しくは2に示すアミノ酸配列、又は該アミノ酸配列の1若 しくは数個のアミノ酸が欠失、 置換若しくは付加されたアミノ carrying out the cloning of the gene which codes alkali cellulase from said microorganisms, it perfected this invention.

[0007]

This invention provides the alkali cellulase gene which codes the amino acid sequence by which the amino acid sequence shown in sequence number 1 or 2, 1 of this amino acid sequence, or some amino acids were delete, substitute or added.

Moreover, this invention provides the alkali cellulase gene which has the base sequence by which the base sequence shown in sequence number 3 or 4, 1 of this base sequence, or some bases were delete, substitute or added. Moreover, this invention provides the recombinant vector containing the above-mentioned alkali cellulase gene, and the transformed body containing this recombinant vector.

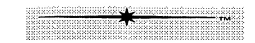
Moreover, this invention cultivates the above-mentioned transformed body.

本発明は、上記の形質転換体を It provides the production of the alkali cellulase 培養することを特徴とするアル characterized by the above-mentioned.

[8000]

[EMBODIMENT OF THE INVENTION]

The gene of this invention has the sequence which codes the amino acid sequence by which the amino acid sequence shown in sequence number 1 or 2, 1 of this amino acid sequence, or some amino acids were delete, substitute or



酸配列をコードする配列を有する。アルカリセルラーゼ活性を 失わない限り、該アミノ酸配列は 中のアミノ酸の欠失、置換又は 付加(以下、置換ということ がある)は特に制限されない。 は特に制限されない。 は熟酵素のアミノ酸配列における アミノ酸がよれるアミノ酸が るアミノ酸が るアミノ酸が なアミノ酸が なアミノ酸が なアミノ酸が なアミノ酸が なアミノでもよい。

[0009]

本発明の配列番号1に示すアル カリセルラーゼ(以下、N13 アミノ酸配列と従来公知のセル 性を比較すると、<u>Bacillus</u> sp. No.1139 株の生産するセルラー ゼ (Fukumori ら、J.Gen. 131, 3339-3345, Microbiol.. 1985) との相同性は81.9% であり、<u>Bacillus</u> sp. KSM-64 株 由来のセルラーゼ(Sumitomo ら Biosci. Biotechnol. Biochem., 56, 872-877, 1992) との相同性は83.6%、 Bacillus sp. KSM-S237 株が生 産するセルラーゼ(特願平11 -013049号) との相同性 は86.7%であり、本発明の 遺伝子からコードされるN13 1 a セルラーゼと最も高い相同 性を示したが、完全に一致する ものではなかった。このことは、

added.

Unless alkali cellulase activity is lost, deletion of the amino acid in this amino acid sequence, substitution, or addition (it may call it variation hereafter) in particular is not limited.

Moreover, as for the amino terminus in the amino acid sequence of the mature enzyme shown in sequence number 1 or 2, one or more amino acids may be added, deleted or replaced.

[0009]

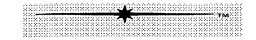
When the homology of the amino acid sequence of alkali cellulase (it shows it as 1 a セルラーゼと表記する) の N131a cellulase hereafter) and the amino acid of conventionally sequence well-known ラーゼのアミノ酸配列との相同 cellulase which are shown in sequence number 1 of this invention is compared, the homology with the cellulase (Fukumori J.Gen.Microbiol., 131, 3339-3345, 1985) which a Bacillus sp.No. 1139 strain produces is 81.9%.

The homology with the cellulase (Japanese Patent Application No. 11-013049) in which 237 strain of Bacillus sp.KSM-S produces the homology with the cellulase (Sumitomo et al., Biosci.Biotechnol.Biochem., 56, 872-877, 1992) derived from Bacillus sp.KSM-64 strain 83.6% is 86.7%.

The N131a cellulase coded from the gene of this invention and the highest homology were shown.

However, it was not what is completely in agreement.

This suggests that N131a cellulase is new alkali



N131aセルラーゼが新規なアルカリセルラーゼであることを示唆するものであり、従って配列番号1に示したアミノ酸配列と最大87%以上の相同性を有するセルラーゼは本発明に含まれる。

cellulase.

Therefore, the amino acid sequence shown in sequence number 1 and the cellulase which has a maximum of 87 % or more homology are contained in this invention.

[0010]

すアルカリセルラーゼ(以下、 N131bセルラーゼと表記す る)のアミノ酸配列と従来公知 のセルラーゼのアミノ酸配列と N131aセルラーゼとの相同 性は83.6%、Bacillus sp. No.1139 株の生産するセルラー ゼとの相同性は88.0%、 Bacillussp. KSM-64 株由来のセ ルラーゼとの相同性は90. 9%であった。さらに、Bacillus sp. KSM-S237 株が生産するセ ルラーゼとの相同性が94. 7%と本発明の遺伝子からコー ドされるN131bと最も高い 相同性を示した。このことは、 するものではなく、新規な酵素 であることを示唆するものであ り、従って配列番号2に示した 本発明に含まれる。尚、相同性 the の検索はGENENTYX-C

[0010]

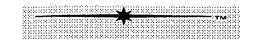
次に、本発明の配列番号2に示 Next, when the homology of the amino acid sequence of alkali cellulase (it shows it as N131b cellulase hereafter) and the amino acid conventionally sequence of well-known cellulase which are shown in sequence number の相同性を比較すると、上記の 2 of this invention was compared, the homology with the cellulase derived from Bacillussp. KSM-64 strain of the homology with the cellulase which, as for the homology with the above-mentioned N131a cellulase, a Bacillus sp.No. 1139 strain produces 83.6% was 90.9% 88.0%.

Furthermore, the homology with the cellulase which 237 strain of Bacillus sp.KSM-S produces showed 94.7%, N131b coded from the gene of this invention, and the highest homology.

ドされるN 1 3 1 b と最も高い N131b cellulase of conventionally well-known 相同性を示した。このことは、 cellulase does not correspond completely, and N 1 3 1 b セルラーゼが従来公 this suggests that it is a new enzyme.

知のセルラーゼとは完全に一致 Therefore, the amino acid sequence shown in するものではなく、新規な酵素 sequence number 2 and the cellulase which であることを示唆するものであ has a maximum of 95 % or more homology are り、従って配列番号2に示した contained in this invention.

アミノ酸配列と最大95%以上 In addition, it performed the search of homology の相同性を有するセルラーゼは with the maximum matching method which used 本発明に含まれる。尚、相同性 the GENENTYX-CD bio-data software の検索はGENENTYX-C [software-development company make and



Dバイオデータソフトウェア ver.36]. 「ソフトウェア開発社製、ve r. 36]を用いたマキシマム マッチング法にて行った。

[0011]

本発明のアルカリセルラーゼ遺 伝子は、配列番号1若しくは2 が欠失、置換若しくは付加され desirable. た塩基配列を有するものが好ま しい。

[0012]

を有するバチルス エスピー mycological characteristics. ーン化することができる。

〔バチルス エスピー KSM Bacillus sp -N131株の菌学的性質]

A. 形態学的性質;

- (a) 細胞の形及び大きさ:桿 to 7.2 micrometer) 菌(0.6~0.8×2.8~ 7. $2 \mu m$
- (b) 多形性:無し
- (c)運動性:有り
- 1. 0×1 . $0 \sim 1$. $8 \mu \text{ m}$

[0011]

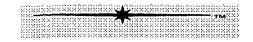
The alkali cellulase gene of this invention should just code the amino acid sequence に示すアミノ酸配列又はその変 shown in sequence number 1 or 2, or its variant. 異体をコードするものであれば However, what has the base sequence by よいが、配列番号3若しくは4 which the base sequence shown by sequence で示される塩基配列、又は該塩 number 3 or 4, 1 of this base sequence, or 基配列の1若しくは数個の塩基 some bases were delete, substitute or added is

[0012]

本発明のアルカリセルラーゼ遺 The alkali cellulase genes of this invention are 伝子は、バチルス属に属する微 the microorganisms belonging to the Bacillus, 生物、例えば下記の菌学的性質 for example, Bacillus sp which has the following It can carry out KSM-N131株等からクロ a cloning from 131 strain of KSM-N etc.

> [Mycological characteristics of 131 strain of KSM-N]

- Morphological characteristic;
- (a) Form and size of cell: Bacillus (0.6-0.8*2.8
- (b) Polymorphism: nothing
- (c) Manoeuverability: be.
- (d) 胞子の形、大きさ、位置、 (d) Form of spore, size, position, existence of 膨潤の有無:楕円形、0.7~ swelling : ellipse form, 0.7-1.0*1.0 to 1.8 micrometer, center semi- end, those with



中央準端、膨潤有り

(e) グラム染色:陽性

swelling

(e) Gram's stain: positive

(f) 抗酸性: 陰性

(f) Acid-fastness: negativity

[0013]

B. 培養学的性質;

- (a) 一般細菌用液体培地(p
- H5.7、培地1):生育せず
- (b) 一般細菌用液体培地 (p
- H 6. 8、培地 1): 生育せず
- (c) 一般細菌用寒天培地 (p
- H6. 5、培地2): 生育せず

[0013]

- B. Culture study characteristic;
- (a) Broth for standard bacteria (pH5.7, medium
- 1): don't grow.
- (b) Broth for standard bacteria (pH6.8, medium
- 1): don't grow.
- (c) Agar for standard bacteria (pH6.5, medium
- 2): don't grow.
- H8. 5、培地2): 生育する
- (d) 一般細菌用寒天培地 (p (d) Agar for standard bacteria (pH8.5, medium
 - 2): grow.

[0014]

C. 生理学的性質;

- 陽性
- (b) 脱窒反応(培地3): 陰性 negativity
- 性

[0014]

- C. Physiological characteristic;
- (a) 硝酸塩の還元(培地3): (a) Reduction of nitrate (medium 3): positive
 - (b) Denitrification reaction (medium 3) :

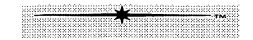
(e) Formation of hydrogen sulfide (medium 6):

(f) Hydrolysis of starch (medium 7): positive

- (c) VPテスト (培地4):陰 (c) VP test (medium 4): negativity
- (d) インドールの生成(培地 (d) Formation of indole (medium 5): negativity
- 5):陰性
- (e) 硫化水素の生成(培地 negativity
- 6):陰性
- (f) デンプンの加水分解(培 (g) Hydrolysis of casein (medium 8): negativity
- 地7):陽性
 - (g) カゼインの加水分解(培
- 地8):陰性

- (h)ゼラチンの液化(培地 (h) Liquefying of gelatin (medium 9): positive
- 9):陽性

- Utilization of 1 citric acid (medium 10) :

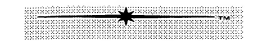


- (i) クエン酸の利用(培地1 negativity
- 0):陰性
- (i) カタラーゼ:陽性
- (k) オキシダーゼ(培地1
- 1):陽性
- 23 38%
- (m) 生育の p H範囲 (培地 1
- 適範囲:pH9-9.5
- (培地14):嫌気条件下で微弱 feeble, grow.
- (o) グルコースからのガス産 negativity 生(培地15):陰性
- 存在下で生育する。
- (q) 馬尿酸の加水分解(培地 negativity
- 17):陰性

だが生育する。

- リルーβーDーグルクロニド (medium 18): negativity
- 8):陰性
- クトース、シュークロース、マ ンノース、マルトース、ラクト ース、トレハロース、フラクト ース、メリビオース、リボース、 サリシン、グリセロール、ソル ビトール等を炭素源として生育

- (i) Catalase : positive
- (k) Oxidase (medium 11): positive
- (1) 生育の温度範囲(培地1 (I) Temperature range of growth (medium 12):
- 2):13-42℃、至適範囲: 13 to 42 degree C, optimum range:23-38 degree C
- (m) The pH range of growth (medium 13): 3): p H 7. 6-10. 5、至 pH7.6-10.5, optimum range:pH9-9.5
 - (n) Influence of the oxygen in growth (medium
- (n) 生育における酸素の影響 14): on anaerobic conditions, although it is
 - (o) Gas production from glucose (medium 15):
- (p) 塩化ナトリウム耐性(培 (p) Sodium chloride resistance (medium 16): 地16):10%塩化ナトリウム grow in a sodium chloride presence 10%.
 - (q) Hydrolysis of hippuric acid (medium 17):
- Hydrolysis of 4-methyl (r) (r) 4ーメチルウンベリフェ umbelliferyl-(beta)-D-glucuronide (MUG)
- (MUG) の加水分解(培地1 (s) Utility of saccharide (medium 19): it can grow the glucose, the arabinose, the xylose, a (s) 糖の利用性(培地19): mannitol, the galactose, sucrose, the mannose, グルコース、アラビノース、キ the maltose, a lactose, a trehalose, a fructose, シロース、マンニトール、ガラ the melibiose, the ribose, the salicin, a glycerol, sorbitol, etc. as a source of a carbon.
 - It cannot utilize a rhamnose and an inositol as a source of a carbon.



可能である。ラムノース、イノ シトールを炭素源として利用で きない。

[0015]

にてpHを調整

トリウムにてpHを調整

0. 1重量% (別滅菌)

ス0.5重量% (別滅菌)、炭酸 carbonate ナトリウム 0.2重量% (別滅 菌)

指示量、炭酸ナトリウム 0.1 重量%(別滅菌)、インドール産 生試験用濾紙 (日水製薬) 培地6: TS I 寒天培地(栄研 化学) 指示量、炭酸ナトリウム 0. 1 重量% (別滅菌) 培地7:バクトペプトン1.5 重量%、酵母エキス0.5重 量%、可溶性デンプン2.0重 量%、リン酸1水素カリウム0. 1重量%、硫酸マグネシウム7 水塩0.02重量%、寒天1. 5重量%、炭酸ナトリウム0.

[0015]

培地1:ニュートリエントブロ Medium 1: The amount of nutrient-broth (Difco) ス(ディフコ)指示量、希塩酸 commands and the diluted hydrochloric acid adjust pH.

培地2:ニュートリエントアガ Medium 2: Adjust pH in the amount of nutrient ー(ディフコ)指示量、炭酸ナ agger (Difco) commands, and the sodium carbonate.

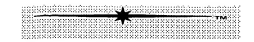
培地3:ニュートリエントブロ Medium 3: 0.8 weight% of nutrient broth, 0.1 ス 0. 8 重量%、硝酸カリウム weight% of potassium nitrate, 0.1 weight% 0. 1 重量%、炭酸ナトリウム (another sterilization) of sodium carbonate

Medium 4: 0.7 weight% (Difco) of bacto 培地4:バクトペプトン(ディ peptone, 0.5 weight% of sodium chloride, 0.5 フコ) 0. 7重量%、塩化ナト weight% (another sterilization) of glucose, 0.2 リウム 0. 5 重量%、グルコー weight% (another sterilization) of sodium

培地5:SIM培地(日水製薬) Medium 5: The amount of SIM medium (Nissui Pharmaceuticals) commands, 0.1 weight% (another sterilization) of sodium carbonate, the filter paper for an indole production test (Nissui Pharmaceuticals)

> Medium 6: The amount of TSI agar (Eiken Chemical) commands, 0.1 weight% (another sterilization) of sodium carbonate

> Medium 7: 1.5 weight% of bacto peptone, 0.5 weight% of yeast extract, 2.0 weight% of soluble starch, 0.1 weight% of phosphoric-acid hydrogen potassium, 0.02 weight% of magnesium-sulfate heptahydride, 1.5 weight% of agar, 0.2 weight% (another sterilization) of



2重量% (別滅菌)

培地8:酵母エキス0.5重 Medium 8:0.5 weight% of yeast extract, 2.0 量%、グルコース2.0重量%、 1水素カリウム0.1重量%、 硫酸マグネシウム7水塩、0. 02重量%、寒天1.5重量%、 炭酸ナトリウム 0.1 重量% (別 carbonate 滅菌)

weight% of glucose, 0.5 weight% of casein, 0.1 カゼイン 0.5 重量%、リン酸 weight% of phosphoric-acid 1 hydrogen potassium, magnesium-sulfate heptahydride, 0.02 weight%, 1.5 weight% of agar, 0.1

sodium carbonate

ス0.8重量%、ゼラチン1. 量%、炭酸ナトリウム 0. 2重 sodium carbonate 量%(別滅菌)

培地10:リン酸1水素アンモ hydrogen ニウム 0. 1 重量%、リン酸 2 水素カリウム 0. 1 重量%、硫 magnesium-sulfate 酸マグネシウム7水塩、0.0 0.2重量%、寒天1.5重量%、 滅菌)

培地11:チトクロムオキシダ ーゼ試験濾紙(日水製薬) ソイ ブロス (BBL) 指示量、 炭酸ナトリウム 0.1 重量%(別 滅菌)

滅菌後に添加し、pHを調整 ー(ディフコ)指示量、炭酸ナ sterilization) of sodium carbonate

培地9:ニュートリエントブロ Medium 9: 0.8 weight% of nutrient broth, 1.2 weight% of gelatin, 0.5 weight% of yeast 2重量%、酵母エキス0.5重 extract, 0.2 weight% (another sterilization) of

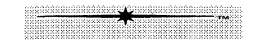
weight% (another sterilization) of sodium

Medium 10: 0.1 weight% of phosphoric-acid 1 ammoniums, 0.1 weight% monobasic potassium phosphate, 0.02 heptahydride, weight%, 0.2 weight% of sodium citrate, 1.5 2重量%、クエン酸ナトリウム weight% of agar, 0.1 weight% (another sterilization) of sodium carbonate

炭酸ナトリウム 0.1 重量% (別 Medium 11: Cytochrome oxidase test filter paper (Nissui Pharmaceuticals)

Medium 12: Tryptocase soy The amount of broth (BBL) commands, 0.1 weight% (another 培地12:トリプティケース sterilization) of sodium carbonate

培地13:トリプティケース Medium 13: Tryptocase soy It adds the ソイ ブロスに炭酸ナトリウム sodium carbonate or the sodium hydroxide to a あるいは水酸化ナトリウムを別 broth after another sterilization, it adjusts pH. Medium 14: The amount of anaerobic agger 培地14:アナエロビックアガ (Difco) commands, 0.2 weight% (another



トリウム0.2重量% (別滅菌) 培地15:バクトペプトン1. 0重量%、塩化ナトリウム0. 5重量%、グルコース1.0重 量%、フェノールレッド0.0 02重量%、水酸化ナトリウム にてpHを調整

培地16:バクトトリプトン(デ ィフコ) 0.5重量%、酵母エ キス1.5 重量%、リン酸1水 素カリウム0.3重量%、寒天 2. 0重量%、グルコース2. 0 重量% (別滅菌)、塩化ナトリ ウム0-16重量%、炭酸ナト リウム 0.5 重量% (別滅菌)

Medium 15: Adjust pH in 1.0 weight% of bacto peptone, 0.5 weight% of sodium chloride, 1.0 weight% of glucose, 0.002 weight% of phenol red, and the sodium hydroxide.

Medium 16: 0.5 weight% (Difco) bactotryptons, 1.5 weight% of yeast extract, 0.3 weight% of phosphoric-acid 1 hydrogen potassium, 2.0 weight% of agar, 2.0 weight% (another sterilization) of glucose, 0 to 16 weight% of sodium chloride, 0.5 weight% (another sterilization) of sodium carbonate

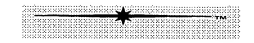
培地17:バクトトリプトン1. 0重量%、肉エキス(ディフコ) 0. 3重量%、酵母エキス0. 量%、リン酸1水素ナトリウム 0.5重量%、馬尿酸1.0重 量%、炭酸ナトリウム 1. 0重 carbonate 量% (別滅菌)

培地18:バクトトリプトース ステイン塩酸塩0.1重量%、 寒天1. 5重量%、MUG10 0 p p m (濾過滅菌)、炭酸ナト リウム 0. 3重量% (別滅菌) 培地19:硝酸カリウム0.2 重量%、リン酸1水素ナトリウ ム0.2重量%、塩化ナトリウ ム0.5重量%、硫酸マグネシ ウム7水塩0.005重量%、

Medium 17: 1.0 weight% of bactotryptons, 0.3 weight% (Difco) of meat extracts, 0.1 weight% of yeast extract, 0.1 weight% of glucose, 0.5 1重量%、グルコース 0. 1重 weight% of phosphoric-acid 1 hydrogen sodium, 1.0 weight% of hippuric acid, 1.0 weight% (another sterilization) of sodium

Medium 18: 2.0 weight% (Difco) of bacto tryptose, 0.5 weight% of sodium chloride, 0.1 (ディフコ) 2. 0 重量%、塩 weight% of cystein hydrochloride, 1.5 weight% 化ナトリウム 0. 5 重量%、シ of agar, MUG100 ppm (filtration sterilization), 0.3 weight% (another sterilization) of sodium carbonate

> Medium 19: 0.2 weight% of potassium nitrate, 0.2 weight% of phosphoric-acid 1 hydrogen sodium, 0.5 weight% of sodium chloride, 0.005 weight% of magnesium-sulfate heptahydride, 0.2 volume % of trace amount metal <mixed-liquid SP> *</SP>, 0.2 volume % of <mixed-liquid SP> **</SP>, the vitamin



緩衝液(pH10)0.1M、 寒天0.3重量%(別滅菌)、糖 類1.0重量%(濾過滅菌) * 、 * * ; Nielsen ら、 141, Microbiology, 1745-1761(1995)に準ずる。

微量金属混液 0. 2容量%、ビ carbonic acid buffer (pH10) 0.1M, 0.3 weight% タミン混液**0.2容量%、炭酸 (another sterilization) of agar, 1.0 weight% (filtration sterilization) of sugars

> It applies to Nielsen et al., Microbiology, 141,1745-1761 (1995).

[0016]

性培地に生育しない好アルカリ 性細菌であり、且つグラム陽性、 カタラーゼ陽性の有胞子桿菌で あることから、好アルカリ性バ た。そこで本菌株の形態学、生 理学的性質について、Nielsen らが新たに分類した好アルカリ 性バチルス属細菌の記載 this-microbe (Microbiology 、 141 、 検討した結果、本菌株はバチル に近縁な菌種であると考えられ た。しかし、その性質は既知の バチルス シュウドアルカロフ ィルスと完全には一致せず、他 known のバチルス属菌の諸性質とも一 致しないため、新規なバチルス 属細菌として本菌株を工業技術 院生命工学研究所へ、バチルス (FERMP-17475) と して寄託した。

[0016]

以上、KSM-N131株は中 As mentioned above, 131 strain of KSM-N is alkalophilic bacteriums which it does not grow to a neutral medium.

And since it was a gram-positive and catalase electropositive owner spore Bacillus, it was チルス属細菌であると判断され judged that they were alkali-loving Bacillus bacteria.

Then, Nielsen and others did comparison examination about the morphology strain, and а physiological characteristic according to publication (1745-1745-1761、1995) に準じ比較 Microbiology, 141, 1761, 1995) of the newly categorized alkali-loving Bacillus bacteria.

ス シュウドアルカロフィルス As a result, it was thought that this-microbe strain was a microbial species with close relation to a Bacillus shoed alcalophilus.

However, since it is not in agreement with a Bacillus shoed alcalophilus and completeness and in agreement with the characteristics of several of another Bacillus genus, the characteristic is Bacillus sp to an institute-of-technology biotechnology research エスピー KSM-N131株 laboratory about this-microbe strain as new Bacillus bacteria. It deposited as 131 strain (FERMP-17475) of KSM-N.



[0017]

知の手段、例えばショットガン 法、PCR法を用いて行うこと PCR method. ができる。

[0018]

また、本発明のアルカリセルラ ーゼ遺伝子を含む組換えベクタ ーを作製するには、宿主内で複 製維持が可能で、該酵素を安定 伝子を安定に保持できるベクタ ーにアルカリセルラーゼ遺伝子 を組込めばよい。かかるベクタ ーとしては大腸菌を宿主とする 場合、pUC18、pBR32 2、pHY300PLK等が挙 げられ、枯草菌を宿主にする場 合、pUB110、pHSP6 (Sumitomo 5 Biosci. Biotechnol. Biochem., 59, 2172-2175, 1995) \ p H Y 3 0 0 P L K 等が挙げられる。

[0019]

かくして得られた組換えベクタ ーを用いて宿主菌を形質転換す るには、プロトプラスト法、コ ンピテントセル法、エレクトロ ポレーション法等を用いて行う ことができる。宿主菌としては 特に制限されないが、Bacillus 属(枯草菌)等のグラム陽性菌:

[0017]

上記のKSM-N131株から As the cloning method of the alkali cellulase のアルカリセルラーゼ遺伝子の gene from 131 strain of above-mentioned クローニング方法としては、既 KSM-N, they are known means, for example, it can carry out using the shotgun method and

[0018]

Moreover, what is necessary is for duplication maintenance to be possible within the host, and to be able to let this enzyme express stably and just to integrate an alkali cellulase gene in the に発現させることができ、該遺 vector which can maintain this gene stably, in order to produce the recombinant vector containing the alkali cellulase gene of this invention.

> When making an Escherichia coli into the host this vector, pUC18, pBR322, pHY300PLK etc. are mentioned, when making the Bacillus subtilis into the host, pUB110, pHSP64 (Sumitomo et al.. Biosci.Biotechnol.Biochem., 59, 2172-2175, 1995), and pHY300PLK etc. are mentioned.

[0019]

In order to transform a host microbe using the recombinant vector obtained by the thing which write, and to do, it can carry out using the protoplast method, the competent cell method, the electroporation method, etc.

It does not limit particularly as a host microbe. However, gram positive bacteria, such as a Bacillus genus (Bacillus subtilis);



グラム陰性菌; Streptomyces 属 coli (Escherichia coli); (放線菌)、Saccharomyces 属 Fungi, (酵母)、<u>Aspergillus</u>属(カビ) 等の真菌が挙げられる。

[0020]

得られた形質転換体を培養し、 当該培養液からアルカリセルラ ーゼを採取することにより、ア 質転換株が資化しうる炭素源、 含む培地を用いて適当な条件下 によって酵素の採取、精製を行 い、凍結乾燥、噴霧乾燥、結晶 化等により、所望の酵素形態と することができる。

[0021]

【実施例】

生産菌のスクリーニング)

日本各地の土壌を滅菌水に懸濁 したものを80℃、30分間熱 天平板培地に塗布した [2.0] 重量%カルボキシメチルセルロ ース(A 1 0 M C; 日本製紙社 compositions (A10MC;)

Escherichia coli (大腸菌) 等の Gram negative bacteria, such as Escherichia

such as a Streptomyces genus (actinomycetes), a Saccharomyces genus (yeast), and an Aspergillus genus (fungi), are mentioned.

[0020]

It cultivates the obtained transformed body, by collecting alkali cellulase from said culture medium, it can obtain alkali cellulase.

ルカリセルラーゼを得ることが What is sufficient is just to perform a culture on できる。培養は、宿主菌又は形 suitable conditions using the medium containing the source of a carbon which a host microbe or 窒素源、金属塩、ビタミン等を the transformant can utilize, the source of nitrogen, a metallic salt, a vitamin, etc.

で行なえばよい。かくして得ら From the culture medium obtained by the thing れた培養液から、一般的な方法 which write, and to do, it can perform collection of an enzyme, and purification by the general method, and can consider it as the desired enzyme form according to freeze-dried, spray drying, crystallization, etc.

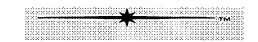
[0021]

[EXAMPLES]

実施例1 (アルカリセルラーゼ Example 1 (screening of an alkali cellulase producing microbe)

It heat-processes 80 degrees C of things which suspended the soil of every place of Japan in 処理し、以下の組成を有する寒 the sterilized water for 30 minutes, [2.0-weight%] carboxymethylcellulose applied to the agar planar medium which has the following

製)、1.0重量%肉エキス(オ The Nippon Paper Industries make, the キソイド社製)、1.0重量%バ 1.0-weight% meat extract (oxo id shrine make),



1. 0 重量%塩化ナトリウム、 0. 1 重量%リン酸 2 水素カリ ウム、0.5重量%炭酸ナトリ ウム(別滅菌)、0.005重量% トリパンブルー (別滅菌)]。3 0℃の培養器で3日間静置培養 し、生育した菌の周辺にカルボ キシメチルセルロースの分解に 伴う溶解斑が検出されたものに ついて選抜し、シングルコロニ 一化を繰り返した。これらの菌 株を、2.0重量%ポリペプト ンS (日本製薬社製)、1.0重 量%魚肉エキス(和光純薬社 製)、0.15重量%リン酸1水 素カリウム、0.1 重量%酵母 エキス (ディフコ社製)、0.0 7重量%硫酸マグネシウム7水 Purechemical 塩、0.1重量%カルボキシメ 量%炭酸ナトリウム (別滅菌) から成る液体培地を用い、3 0℃、3日間振盪培養した。ア ルカリセルラーゼを生産してい る菌株を選択し、とりわけ高ア ルカリ性域で強力な活性を示し たセルラーゼ生産菌としてバチ ルス エスピー KSM-N1 31株を取得した。

[0022]

KSM-N131株のゲノムD 131 strain of Bacillus sp NAの調製)

クトペプトン(ディフコ社製)、 the 1.0-weight% bacto peptone (made by a Difco company), 1.0-weight% sodium chloride, 0.1-weight% monobasic potassium phosphate, the 0.5-weight% sodium carbonate (another sterilization), and the 0.005-weight% trypan blue (another sterilization)].

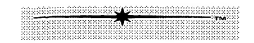
> It carries out stationary culture for three days by a 30-degree C incubator, it selects about that from which the melting spots accompanying a degradation of carboxymethylcellulose were detected around the grown microbe, it repeated single colony-ization.

It carried out the shaking culture of the 30 degrees C of these strains for three days using the broth which constitutes of the 2.0-weight% polypeptone S (made by NIHON PHARMACEUTICAL CO., LTD.), the 1.0-weight% fish-meat extract (made by a Wako KK), 0.15-weight% phosphoric-acid 1 hydrogen potassium, the チルセルロース及び 0.5 重 0.1-weight% yeast extract (made by a Difco company), 0.07-weight% magnesium-sulfate heptahydride, 0.1-weight% carboxymethylcellulose, and the 0.5-weight% sodium carbonate (another sterilization).

> It chooses the strain which produces alkali cellulase, it is Bacillus sp as a cellulase producing microbe which showed activity especially powerful in a high alkaline region. It acquired 131 strain of KSM-N.

[0022]

実施例2(バチルス エスピー Example 2 (manufacture of the genome DNA of KSM-N) Bacillus sp Using the medium which バチルス エスピー KSMー constitutes of the 2.0-weight% polypeptone S,



量%カルボキシセルロース(A fish-meat 10MC)、0.1重量%酵母工 キス、1重量%魚肉エキス、0. 15 重量%リン酸1水素カリウ ム、0.07重量%硫酸マグネ シウム 7 水塩、 0. 5 重量%グ ルタミン酸ナトリウム(別滅菌) 及び0.5重量%炭酸ナトリウ ム(別滅菌)から成る培地を用 い、30℃、40時間振盪(1 25 r p m) して行った。得ら れた培養液約300mLから遠 心分離(12000×g、15 分、5℃) により菌体を回収し、 この菌体から斉藤・三浦の方法 によりゲノムDNAを調製し た。

N 1 3 1 株の培養は、2.0重 the 0.1-weight% carboxy cellulose (A10MC), 量%ポリペプトンS、0. 1重 the 0.1-weight% yeast extract, the 1-weight% extract. 0.15-weight% phosphoric-acid 1 hydrogen potassium, 0.07-weight% magnesium-sulfate heptahydride, the 0.5-weight% sodium glutamate (another sterilization), and the 0.5-weight% sodium carbonate (another sterilization), 30 degrees C, it shook the culture of 131 strain of KSM-N for 40 hours (125 rpm), and performed it.

> Centrifugation (12000*g, 15 minutes, 5 degrees C) recovers a microbial cell from about 300 mL of obtained culture mediums, it prepared genome DNA by the method of Saito and a Miura from this microbial cell.

[0023]

ゼ遺伝子断片のクローニング) バチルス エスピー KSM-N131株の培養上清から精製 したセルラーゼのアミノ末端配 列を15番目まで決定した結 果、Glu-Gly-Asn- As Thr-Arg-Glu-Asp-A s n-P h e-A s p-His-Leu-Leu-Gly-Asnであった。この配列 は、バチルス エスピー KS M-S237株やバチルス スピー KSM-64株の生産 するアルカリセルラーゼのアミ

[0023]

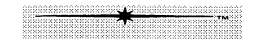
実施例3(N131aセルラー Example 3 (cloning of an N131a cellulase gene fragment)

> It decided the amino-terminus Bacillus sp sequence of the cellulase purified from the culture supernatant of 131 strain of KSM-N to the 15th.

> а result, it was Glu-Gly-Asn-Thr-Arg-Glu-Asp-Asn-Phe-Asp-His -Leu-Leu-Gly-Asn.

237 strain of This sequence is Bacillus sp. KSM-S, and Bacillus sp Amino-terminus sequence

Glu-Gly-Asn-Thr-Arg-Glu-Asp-Asn-Phe-Lys-His -Leu-Leu-Gly-Asn of the KSM-64 strain alkali cellulase to produce and extremely high



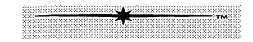
ノ末端配列G1u-G1y-A sn-Thr-Arg-Glu-A s p - A s n - P h e - Lys-His- Leu-Le u-Gly-Asnと極めて高 い相同性を示した。そこで中間 のアミノ酸配列も相同性が高い 可能性があると予想し、S23 7セルラーゼのアミノ末端及び 中間アミノ酸配列を基にプライ マー1 (配列番号5) 及びプラ イマー2 (配列番号6) を合成 し、これらを用いてN131a セルラーゼをコードする遺伝子 の増幅をPCR反応により試み た。すなわち、バチルス エス ピー KSM-N131株ゲノ ム溶液 1 μ L (100 n g)、プ ライマー1及び2各20μL (1 μ M)、P C R 用緩衝液 1 0 μL , 2. 5 mMdNTP>クス8 μ L、Pyrobest DNAポリメラーゼ(タカラ社 製) 0 . 5 μ L (2 . 5 単位)、 及び脱イオン水 4 0 μ Lを混合 し、サーマルサイクラー480 (パーキンエルマー社製) にて 94℃、2分間の熱変性後、9 4℃で1分間、55℃で1分間、 72℃で2分間を1サイクルと し、30サイクルの反応条件で DNAの増幅を行った。得られ たPCR産物(約1kb)をG DNA and FX PCR Gel Band Purif ication Kit (7r

homology were shown.

Then, it also anticipates a middle amino acid sequence that homology may be high, it compounds primer 1 (sequence number 5) and primer 2 (sequence number 6) based on the amino terminus and middle amino acid sequence of S237 cellulase, it tried amplification of the gene which codes N131a cellulase using these according to PCR reaction.

Namely, Bacillus sp It mixes 131 strain of KSM-N genome solution 1 micronL (100 ng), primer 1 and 220 micronL each (1 micronM), buffer 10 micronL for PCR, 2.5 mMdNTP mix 8 micronL, PyrobestDNA polymerase (made by Takara company) 0.5 micronL (2.5 unit), and deionized-water 40 micronL, after 94 degrees C and the thermal denaturation for 2 minutes, by 94 degrees C, it makes for 1 minute at 55 degrees C for 1 minute, and makes for 2 minutes into 1 cycle at 72 degrees C at thermal cycler 480 (made by Perkin-Elmer corporation), it performed amplification of DNA on 30-cycle reaction conditions.

GFX PCR DNA and Gel Band Purification Kit (made by Pharmacia K.K.) purifies the acquired PCR production (about 1 kb), it decided the base sequence of the obtained DNA fragment using DNA Sequencing Kit (made by an applied bio-system company), and a 377DNA sequencer (made by a Perkin-Elmer bio-system company).



ルマシア社製)により精製し、 得られたDNA断片の塩基配列 **EDNA** Sequencin Kit(アプライドバイオ システム社製)及び377DN Aシークエンサー(パーキンエ ルマーバイオシステム社製)を 用いて決定した。

[0024]

るクローニング)

のであったため、インバースP CR法により全遺伝子の取得を 試みた。すなわち、バチルス エ スピー KSM-N131株ゲ ノム溶液10μL (8μg)、P 水34μL及びEcoRI1μ L(10単位)を混合し、37℃、 た。得られたゲノム分解産物を 精製後、Ligation K it Ver. 2 (タカラ社製) を用いて自己閉環した(16℃、 2時間)。自己閉環したDNAを PCR method. 鋳型として用いた。PCR反応 は、自己閉環溶液1 µ L、プラ イマー3(配列番号7)及びプ ライマー4(配列番号8)各2

[0024]

実施例4(N131aセルラー Example 4 (cloning by the genome PCR ゼ遺伝子のゲノムPCR法によ method of an N131a cellulase gene)

Since the N131a cellulase gene decided in 実施例3で決定したN131a Example 3 was imperfect, it tried acquisition of セルラーゼ遺伝子は不完全なも all genes by Inverse PCR method.

Namely, Bacillus sp It mixes 131 strain of KSM-N genome solution 10 micronL (8 microgram), buffer 5 micronL for PCR, deionized-water 34 micronL, and EcoRI1 micronL (10 unit), 37 degrees C carried out CR用緩衝液 5 μ L、脱イオン restriction enzyme treatment for 2 hours and 30 minutes.

It carried out the self-ring closure after purifying 2時間30分間制限酵素処理し the acquired genome cleavage product using Ligation Kit Ver.2 (made by the Takara company) (16 degrees C, 2 hours).

> It purifies DNA which carried out the self-ring closure, it used as a casting mould of Inverse

精製し、インバースPCR法の PCR reaction is self-ring-closure solution 1 micronL, primer 3 (sequence number 7), and primer 4(sequence number 8) 20 micronL each (1 micronM), buffer 10 micronL for PCR, 2.5 mMdNTP mix 8 micronL, after mixing Pyrobest $0 \mu L (1 \mu M)$ 、PCR用緩衝 DNA-polymerase 0.5 micronL (2.5 unit) and 液 1 0 μ L 、 2 . 5 mM d N T deionized-water 40.5 micronL, after 94 degrees



 $P \in \mathcal{V} \setminus \mathcal{A} \setminus \mathcal{B} \setminus \mathcal{A} \setminus \mathcal{A} \cup \mathcal{A$ est DNAポリメラーゼ 0. 5 μ L (2. 5 単位)、及び 脱イオン水40.5μLを混合 した後、94℃、2分間の熱変 性後、94℃で1分間、55℃ で1分間、72℃で3分間を1 サイクルとし、30サイクル行 った。増幅したDNA断片(約 4 k b) を精製し、このうち約 2 k b の塩基配列を決定した。 この段階で完全なN131aセ ルラーゼ遺伝子及びその上流約 500bの配列は決定されたの で、次に構造遺伝子下流の塩基 配列決定を進め、下流約200 bの塩基配列を決定した。得ら れた塩基配列からセルラーゼ遺 伝子の上流領域並びに下流領域 の塩基配列を基に、プライマー 5 (配列番号9) 及びプライマ 一6(配列番号10)を合成し、 バチルス エスピー N131 株のゲノムからPCR法により N131aセルラーゼ遺伝子を 増幅した。得られた遺伝子の塩 基配列を決定し、アミノ酸配列 を推定した (配列番号1及び 3)。

C and the thermal denaturation for 2 minutes, by 94 degrees C, it makes for 1 minute at 55 degrees C for 1 minute, and makes for 3 minutes into 1 cycle at 72 degrees C, it performed 30 cycles.

It purifies the amplified DNA fragment (about 4 kb(s)), among these, it decided the base sequence of about 2 kb(s).

Since, the sequence of a perfect N131a cellulase gene and its upper approximately 500b was decided in this phase, next, it advanced the base-sequence decision of a structural-gene downstream, and decided the base sequence of down-stream approximately 200b.

Based on the base sequence of the upstream region of a cellulase gene, and a downstream region, it compounds primer 5 (sequence number 9) and primer 6 (sequence number 10) from the obtained base sequence, bacillus sp It amplified the N131a cellulase gene by PCR method from the N131 strain genome.

It decides the base sequence of the obtained gene, it presumed the amino acid sequence (sequence number 1 and 3).

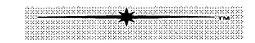
[0025]

産)

[0025]

実施例 5 (形質転換枯草菌によ Example 5 (production of the N131a cellulase るN 1 3 1 a セルラーゼの生 by the transforming Bacillus subtilis)

It connects the gene from the amino-terminus N131aセルラーゼのアミノ side of N131a cellulase to a terminator 末端側からターミネーター下流 downstream with the Sall/BamHI part of a



までの遺伝子をプラスミド(p HSP64) OSalI/Ba mHI部位に連結し、構築した 組換えプラスミドを枯草菌 I S W1214株に導入して形質転 換した。形質転換株を3.0重 量%ポリペプトンS、3.0重 量%マルトース、0.5重量% 魚肉エキス、0.1重量%酵母 エキス、0.1重量%リン酸2 水素カリウム、0.02重量% 硫酸マグネシウム7水塩及びテ トラサイクリン(7.5 μ g/ mL)から成る培地 (PM培地、 pH6. 8) にて30℃、48 時間振盪培養を行った。遠心分 離(8000×g、20分間、 **4℃**) により得られた培養上清 中のセルラーゼの活性は、約2 0000U/Lであった。

plasmid (pHSP64), it introduced the built recombinant plasmid into 1214 strain of Bacillus subtilis ISW, and transformed it.

It performed 30 degrees C and a 48-hour shaking culture in the medium (PM medium, pH6.8) which constitutes the transformant of the 3.0-weight% polypeptone S, the 3.0-weight% maltose, the 0.5-weight% fish-meat extract, the 0.1-weight% yeast extract, 0.1-weight% monobasic potassium phosphate, 0.02-weight% magnesium-sulfate heptahydride, and tetracycline (7.5 microgram/mL).

The activity of the cellulase in the culture supernatant obtained by the centrifugation (for 8000*g and 20 minutes, 4 degrees C) was about 20000 U/L.

[0026]

るクローニング)

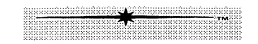
N131aセルラーゼのクロー ニングを行った際に、N131 a セルラーゼの配列と類似した 配列がバチルス エスピー K SM-N131株のゲノム上に 存在する可能性が示唆された。 そこで、N131aセルラーゼ 遺伝子のクローニングの際に用 いた方法と同様に、S237セ ルラーゼのアミノ末端及び中間 アミノ酸配列を基にプライマー

[0026]

実施例6(N131bセルラー Example 6 (cloning by the genome PCR ゼ遺伝子のゲノムPCR法によ method of an N131b cellulase gene)

> The sequence which was similar with the sequence of N131a cellulase when a cloning of N131a cellulase was performed is Bacillus sp. Possibility of existing on the genome of 131 strain of KSM-N was suggested.

> Then, it compounds a primer 7-12 (sequence number 11-16) based on the amino terminus and middle amino acid sequence of S237 cellulase like the method used on the occasion of a cloning of an N131a cellulase gene, it performed amplification of the gene which codes N131b cellulase by PCR method.



7~12(配列番号11~16) を合成し、PCR法によりN1 31bセルラーゼをコードする 遺伝子の増幅を行った。すなわ ち、バチルス エスピー KS M-N131株のゲノム溶液1 組合せを各10μL(0.3μ M)、PCR用緩衝液10μL、 2. 5mMdNTPミックス8 μ L、脱イオン水60μ L及び PwoDNAポリメラーゼ(ベ ーリンガーマンハイム社製) 1 μ L (5単位)を混合し、94℃、 2分間の熱変性後、94℃で1 分間、55℃で1分間、72℃ で3分間を1サイクルとし、3 0サイクルの反応条件でDNA の増幅を行った。得られたPC R産物をHigh Pure PCR Product P u rification Kit (ベーリンガーマンハイム社製) を用いて精製し、377DNA シークンサーにより塩基配列を それぞれ決定した。得られた遺 伝子断片の塩基配列をS237 セルラーゼ遺伝子と比較する と、N131bセルラーゼのア ミノ末端以降をコードすると考 えらるいくつかの遺伝子断片及 び停止コドンとその下流域と考 えられる遺伝子断片の存在が示 唆された。しかし、完全な塩基 配列は決定されていないこと並 びに開始コドン及びその近傍の

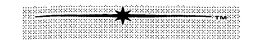
Namely, Bacillus sp It mixes ten micronL(s) each (0.3 micronM), buffer 10 micronL for PCR, 2.5 mMdNTP mix 8 micronL, deionized-water 60 micronL, and PwoDNA polymerase (made by Boehringer-Mannheim company) 1 micronL (5 unit) for the combination of genome solution 1 micronL (70 ng) of 131 strain of KSM-N, and a primer, after 94 degrees C and the thermal denaturation for 2 minutes, by 94 degrees C, it makes for 1 minute at 55 degrees C for 1 minute, and makes for 3 minutes into 1 cycle at 72 degrees C, it performed amplification of DNA on 30-cycle reaction conditions.

It purifies the acquired PCR production using High Pure PCR Product Purification Kit (made by a Boehringer-Mannheim company), 377DNA sequencer- each decided the base sequence.

The presence of the gene fragment considered to be the gene fragment and the stop codon, and its down-stream region of some which are considered to code the amino terminus of N131b cellulase or subsequent ones compared with a S237 cellulase gene in the base sequence of the obtained gene fragment was suggested.

However, the perfect base sequence was not acquired as a gene fragment about not deciding, the initiating codon, and the region of the vicinity.

First, it is Bacillus sp in order to acquire the gene which codes the upstream region from an amino terminus. Various restriction enzymes (Sau 3A, EcoRI, HindIII) degrade genome-DNA 4 microgram of 131 strain of KSM-N, it made what was connected with the cassette using the LA PCR in vitro cloning kit



て取得されていなかった。まず、 アミノ末端より上流の領域をコ ードする遺伝子を取得するため にバチルス エスピー KSM -N131株のゲノムDNA4 μgを各種制限酵素(Sau3 A, <u>Eco</u>RI, <u>Hi</u>ndIII) により分解し、LA PCRイ ンヴィトロクローニングキット (宝酒造)を用いてカセットと連 結したものを鋳型にPCR反応 [プライマー13 (配列番号1 7) 及びプライマー14(配列 番号18)を使用]を行った。 その結果、<u>H i n</u> d Ⅲ により処 理したサンプルについてDNA の増幅が認められ、この増幅断 片(約0.4kb)の塩基配列 を決定した結果、N131bセ ルラーゼのアミノ末端より上流 の領域をコードする遺伝子断片 が確認された。しかし、その解 析を行うと開始コドンから34 塩基下流にアンバーコドン(T GA)が存在することが明らか になった。アンバーコドン(T GA)に関しては、枯草菌にお いて極くまれにトリプトファン をコードするという報告もある ことから(Lovett ら、J. Bacteriol., 173, 1810-1812, 1991)、本遺伝子においてもト リプトファンをコードする可能 性が示唆された。しかし、開始 コドンの上流部にはリボソーム

領域については遺伝子断片とし (Takara Shuzo) for the PCR reaction [primer 13 て取得されていなかった。まず、 (sequence number 17) and primer 14 アミノ末端より上流の領域をコ (sequence number 18) to the casting mould.

As a result, amplification of DNA is observed about the sample treated by HindIII, it decided the base sequence of this amplification fragment (about 0.4 kb(s)).

As a result, the gene fragment which codes the upstream region from the amino terminus of N131b cellulase was checked.

However, when the analysis was conducted, it became clear from the initiating codon that an amber codon (TGA) exists in 34 base downstream.

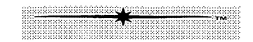
Since there is a report of coding the tryptophan rarely extremely in the Bacillus subtilis about an amber codon (TGA) (Lovett et al., J.Bacteriol., 173, 1810-1812, 1991), possibility of coding the tryptophan was also suggested in this gene.

However, a sequence required for the translation start of a ribosome binding site etc. was not discovered by the upper part of the initiating codon, but it also became clear that many ochre codons (TAA) exist.

Therefore, it was thought that this gene had high possibility of being the false gene which does not express in the cell.

In order to decide the base sequence which codes a perfect N131b cellulase gene, it used primer 15 (sequence number 19) and primer 16 (sequence number 20), and performed PCR reaction.

The base sequence eventually decided and the amino acid sequence presumed were shown in sequence number 2 and sequence number 4.



結合部位などの翻訳開始に必要 な配列が見出されず、オーカー コドン(TAA)がいくつも存 在することも明らかになった。 従って、本遺伝子は細胞内で発 現しない擬似遺伝子である可能 性が高いと考えられた。完全な N131bセルラーゼ遺伝子を コードする塩基配列を決定する ためにプライマー15(配列番 号19)及びプライマー16(配 列番号20)を用いてPCR反 応を行った。最終的に決定され た塩基配列及び推定されるアミ ノ酸配列を配列番号2及び配列 番号4に示した。

[0027]

実施例7(形質転換枯草菌によ 産)

させる目的で、遺伝子の発現に アルカリセルラーゼ遺伝子 (Sumitomo ら、 Biotechnol. Biochem., 56, 827-877、1992)の上流発現領 域を増幅した[プライマー17 (配列番号21) 及びプライマ - 18(配列番号22)を使用]。 得られたN131bセルラーゼ 遺伝子断片と上流発現領域遺伝 子断片を精製し、プライマー1

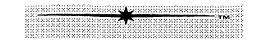
[0027]

Example 7 (production of the N131b cellulase るN131bセルラーゼの生 by the transforming Bacillus subtilis)

It is Bacillus sp as region required for the 細胞内で発現しない可能性のあ expression of a gene in order to produce the るN131b cellulase which may not express in the cell. It is the [primer 17 (sequence number 必要な領域としてバチルス エ 21) and primer 18 (sequence number 22) which スピー KSM-64株由来の amplified the upper expression region of the alkali cellulase gene (Biosci. Biotechnol. Biosci. Sumitomo et al., Biochem., 56,827-877, 1992) derived from KSM-64 strain Use]

> It purifies the N131b cellulase gene fragment and the upper expression region gene fragment which were obtained, recombinant PCR method performed amplification of DNA using primer 16 (sequence number 20) and primer (sequence number 21).

> It purifies the acquired chimera gene, it



6 (配列番号20) 及びプライ マー17(配列番号21)を用 いてリコンビナントPCR法に よりDNAの増幅を行った。取 得したキメラ遺伝子を精製し、 制限酵素Bgl|| びHind||| で処理後、予め同じ制限酵素で 処理しておいたプラスミドpH Y300PLK(ヤクルト本社 製)に連結した。得られた組換 えプラスミドをプロトプラスト 法により枯草菌ISW1214 株に導入し、形質転換を行った。 形質転換株をPM培地(テトラ サイクリンは15μg/mLと した) 中で30℃、72時間振 湿培養した。 遠心分離により得 られた培養上清中のセルラーゼ の活性は、約33000U/L であった。

connected with plasmid pHY300PLK (made by Yakult Honsha) treated beforehand at the same restriction enzyme after treatment by restriction enzyme BgIII HindIII.

It introduces the obtained recombinant plasmid into 1214 strain of Bacillus subtilis ISW by the protoplast method, it performed transforming. It carried out the shaking culture of the 30 degrees C of the transformant for 72 hours in PM medium (it set the tetracycline to 15 microgram(s)/mL).

The activity of the cellulase in the culture supernatant obtained by the centrifugation was about 33000 U/L.

[0028]

[酵素活性測定法] O. 2mL の0.5Mグリシンー水酸化ナ トリウム緩衝液 (pH9.0)、 0.4mLの2.5重量%カル ボキシメチルセルロース(A0 1MC;日本製紙社製)及び0. 3mLの脱イオン水から成る反 応液に、適当に希釈した0.1 mLの酵素液を加えて20分間 反応させた後、1mLのジニト ロサリチル酸試薬(0.5重量% ジニトロサリチル酸、30重 量%ロッシェル塩、1.6重量%

[0028]

[Enzyme active measuring method]

0.2 mL 0.5M glycine- sodium-hydroxide buffer (pH9.0)0.4 mL 2.5-weight% carboxymethylcellulose (A01MC;)

After adding the 0.1 mL enzyme liquid diluted suitably to the reaction mixture which constitutes of the Nippon Paper Industries make and a 0.3 mL deionized water and letting it react to it for 20 minutes, it adds the 1 mL dinitro salicylic-acid reagent (the 0.5-weight% dinitro salicylic acid, the 30-weight% Rochelle salt, the 1.6-weight% sodium-hydroxide aqueous solution), it performed the color development of 水酸化ナトリウム水溶液) を添 the reducing sugar for 5 minutes in boiling



加し、沸水中で5分間還元糖の 発色を行った。氷水中で急冷し、 4mLの脱イオン水を加え、5 して還元糖の生成量を求めた。 尚、ブランクは酵素液を加えず に処理した反応液にジニトロサ ものを用意した。酵素1単位(1 とした。

[0029]

ゼの最適反応 p H)

9)、グリシンー水酸化ナトリウ hydrochloric-acid ム緩衝液 (pH8-11)、リン sodium-hydroxide (pH12-12.5). 適反応 p Hを調べた結果、N 1 9. 5のグリシンー水酸化ナト 速度を示した。また、 p H 7 か (FIG. 1). ら11の間で最大活性の50% 以上の活性を有していた(図 1)。

[0030]

water.

It quenches in ice water, it added the 4 mL deionized water, it measured the absorbence in 3 5 n mにおける吸光度を測定 535 nm, and calculated the produced amount of the reducing sugar.

In addition, a blank adds enzyme liquid, after adding the dinitro salicylic-acid reagent to the リチル酸試薬を加えた後、酵素 reaction mixture treated without adding enzyme 液を添加し、同様に発色させた liquid, it prepared what was developed colors similarly.

U) は、上記反応条件下におい It made 1 unit (1U) of enzymes into the quantity て1分間に1μmolのグルコ which forms the reducing sugar of the glucose ース相当の還元糖を生成する量 of 1 micrometerol in 1 minute on the above-mentioned reaction conditions.

[0029]

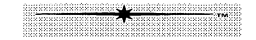
参考例1 (N131a セルラー Reference Example 1 (the optimal reaction pH of N131a cellulase)

クエン酸緩衝液(pH4-7)、 It examined the optimal reaction pH using each リン酸緩衝液 (pH6-8)、ト buffer (100 mM) of citrate buffer solution リスー塩酸緩衝液(pH7- (pH4-7), a phosphate buffer (pH6-8), trisbuffer (pH7-9),glycinebuffer (pH8-11), and 酸-水酸化ナトリウム緩衝液 phosphoric-acid-sodium-hydroxide buffer

衝液(100mM)を用いて最 As a result, N131a cellulase showed the highest reaction rate in the glycine- sodium-hydroxide

Moreover, it had the activity of 50 % or more of リウム緩衝液中で最も高い反応 the maximum activity between 11 from pH7

[0030]



ゼの最適反応 p H)

衝液中で最も高い反応速度を示 was shown. 性を有していた(図2)。

参考例2(N131bセルラー Reference Example 2 (the optimal reaction pH of N131b cellulase)

参考例1と同様にしてN131 It examined the optimal reaction pH of N131b b セルラーゼの最適反応 p Hを cellulase like Reference Example 1.

調べた結果、pH9-9.5の As a result, the highest reaction rate in the グリシン-水酸化ナトリウム緩 glycine- sodium-hydroxide buffer of pH9-9.5

した。また、pH7から11の Moreover, it had the activity of 50 % or more of 間で最大活性の50%以上の活 the maximum activity between 11 from pH7 (FIG. 2).

[0031]

【発明の効果】

伝子を用いれば、衣料用洗剤、 繊維処理剤等として有用なアル に生産することが可能である。

[0031]

[ADVANTAGE OF THE INVENTION]

本発明のアルカリセルラーゼ遺 If the alkali cellulase gene of this invention is used, alkali cellulase useful as the detergent for garments, a fiber processing agent, etc. is カリセルラーゼを単一且つ大量 producible individually and in large quantities.

[0032]

【配列表】

SEQUENCE LISTING

<110> KAO CORPORATION

<120> Gene for

Cellulase

[0032]

[SEQUENCE TABLE]

SEQUENCE LISTING

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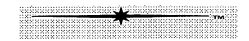
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<400> 1

<212> PRT

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<400> 1



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Leu Val Leu Leu Ser Leu Leu Ala Ala Glu Gly

Phe Pro Thr Ala Leu Ala Ala 20 25

Glu Gly 30

20 Asn Thr Arg Glu Asp Asn Phe Asp His Leu Leu

25 30 Gly Asn Glu Asn Val

Asn Thr Arg Glu Asp Asn Phe Asp His Leu Leu Gly Asn Glu

Asn Val

35 35 40

40 45 45

Lys Arg Pro Ser Glu Ala Gly Ala Lys Arg Pro Ser Glu Ala Gly Ala Leu Gln Leu

Leu Gln Leu Lys Glu Val Asp Lys Glu Val Asp Gly

Gly 50 55

50 55 60

60 Gln Met Thr Leu Val Asp Gln His Gly Glu Lys Ile

Gln Met Thr Leu Val Asp Gln Gln Leu Arg Gly

His Gly Glu Lys Ile Gln Leu Arg

Gly

65 70 65 70

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Trp Phe Pro Glu IIe Leu Asn Leu Asn Asp Asn

Asp Asn 85 90

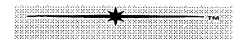
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Ala Tyr Lys Ala Leu Ser Asn Asn Met Ile Arg Leu

Asp Trp Asp Ser Asn Met Ile

Arg Leu

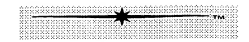


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195 195 200

200 205 205

Lys Ala Val Lys Glu Tyr Ala Asp Lys Ala Val Lys Glu Tyr Ala Asp Pro Ile Val Glu



Pro Ile Val Glu Met Leu Arg Asp Met Leu Arg Asp

210 215 210 215

220 220

Ser Gly Asn Ala Asp Asp Asn Ser Gly Asn Ala Asp Asp Asn Ile Ile Val Gly

Ile Ile Ile Val Gly Ser Pro Asn Ser Pro Asn Trp

Trp

225 230 225 230

235 240 235 240

Ser Gln Arg Pro Asp Leu Ala Ser Gln Arg Pro Asp Leu Ala Ala Asp Asn Pro

Ala Asp Asn Pro Ile Asn Asp His Ile Asn Asp His His

His 245 250

245 255

250 Thr Met Tyr Thr Val His Phe Tyr Ser Gly Ser His

Thr Met Tyr Thr Val His Phe Tyr Ala Ala Ser Thr

Ser Gly Ser His Ala Ala Ser Thr

260 260 265

265 270 270

Glu Ser Tyr Pro Pro Glu Thr Pro Glu Ser Tyr Pro Pro Glu Thr Pro Asn Ser Glu

Asn Ser Glu Arg Gly Asn Val Arg Gly Asn Val Met

Met 275 280

275 285

280 285 Ser Asn Thr Arg Tyr Ala Leu Glu Asn Gly Val Ala

Ser Asn Thr Arg Tyr Ala Leu Val Phe Ala Thr

Glu Asn Gly Val Ala Val Phe Ala

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300 300

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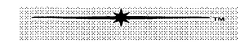
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Leu Asn Glu Asn Asn Ile Ser Trp

325 325 330

330 335 335

Ala Asn Trp Ser Leu Thr Asn Ala Asn Trp Ser Leu Thr Asn Lys Asn Glu Val

Lys Asn Glu Val Ser Gly Ala Ser Gly Ala Phe Thr

Phe Thr 340 345

340 350

345 350 Pro Phe Glu Leu Gly Lys Ser Asn Ala Thr Ser

Pro Phe Glu Leu Gly Lys Ser Leu Asp Pro Gly Pro

Asn Ala Thr Ser Leu Asp Pro

Gly Pro

355 355 360

360 365 365

Asp Gln Val Trp Ala Pro Glu Glu Asp Gln Val Trp Ala Pro Glu Glu Leu Ser Leu

Leu Ser Leu Ser Gly Glu Tyr Val Ser Gly Glu Tyr Val

370 375 370 375

380 380

Arg Ala Arg Ile Lys Gly Ala Lys Arg Ala Arg Ile Lys Gly Ala Lys Tyr Glu Pro Ile

Tyr Glu Pro lle Asp Arg Thr Arg Asp Arg Thr Arg

385 390 385 390

395 400 395 400

Tyr Thr Lys Val Leu Trp Asp Tyr Thr Lys Val Leu Trp Asp Phe Asn Asp Gly

Phe Asn Asp Gly Thr Lys Gln Thr Lys Gln Gly Phe

Gly Phe 405 410

405 415

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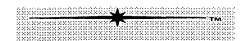
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Asn Lys Glu Ala Ile Glu Val Glu

Asn

420 420 425

425 430 430



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Gly Leu Asn Val Ser Asn Asp Ser Asn Asp Leu

Leu 435 440

435 445

440 445 Ser Asp Gly Asn Phe Trp Ala Asn Val Arg Leu

Ser Asp Gly Asn Phe Trp Ala Ser Ala Asn Gly Trp

Asn Val Arg Leu Ser Ala Asn

Gly Trp

450 455 450 455

460 460

Gly Lys Ser Val Asp Ile Leu Ser Gly Lys Ser Val Asp Ile Leu Ser Ala Glu Lys Leu

Ala Glu Lys Leu Thr Met Asp Thr Met Asp Gly

Gly 465 470

465 470 475 480

475 480 Ile Val Asp Glu Pro Thr Thr Val Ala Ile Ala Ala Ile

Ile Val Asp Glu Pro Thr Thr Val Pro Gln Ser

Ala Ile Ala Ala Ile Pro Gln Ser

485 485 490

490 495 495

Thr Lys His Gly Trp Ala Asn Pro Thr Lys His Gly Trp Ala Asn Pro Glu Arg Ser Val

Glu Arg Ser Val Lys Val Thr Glu Lys Val Thr Glu

500 500 505

505 510 510

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Leu Thr

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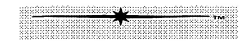
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Leu Lys Asn Ile Gly Phe Asp Gly Phe Asp Asp

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530 535 540

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Glu Asn Asn Met Asn Asn Val Gly Thr Glu Ala Ile Ile Leu Phe Val Gly Thr Glu Ala

545 550 545 550

555 560 555 560

Ala Asp Val Ile Tyr Leu Asp Asn Ala Asp Val Ile Tyr Leu Asp Asn Ile Lys Val Thr

Ile Lys Val Thr Gly Lys Ile Val Gly Lys Ile Val

565 565 570

570 575 575

Glu lle Pro Val Val His Ser Pro Glu lle Pro Val Val His Ser Pro Lys Gly Asp Ala

Lys Gly Asp Ala Ala Leu Pro Ala Leu Pro Ser

Ser

580 580 585

585 590 590

Asn Phe Glu Asp Gly Thr Arg Asn Phe Glu Asp Gly Thr Arg Gln Gly Trp Asp

Gln Gly Trp Asp Trp Ala Gly Glu Trp Ala Gly Glu Ser

Ser 595 600

595 605

600 Gly Val Lys Thr Ala Leu Thr Ile Glu Glu Ala Asn

Gly Val Lys Thr Ala Leu Thr Ile Gly Ser Gln Ala

Glu Glu Ala Asn Gly Ser Gln Ala

610 615 610 615

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Pro Glu Val Lys Pro Ser Asp Pro Ser Asp Asn Trp

Asn Trp 625 630

625 630 635 640

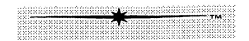
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Phe His Lys Asp Asn Leu Val

Arg Gly

645 645 650



650 655 655

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Asp Phe Tyr IIe Asp Pro Ala Arg Asp Pro Ala Arg Ala

Ala 660 665

660 670

665 670 Thr Glu Gly Ala Met Asn Ile Asn Leu Val Phe

Thr Glu Gly Ala Met Asn Ile Asn Gln Pro Pro Ala Asn

Leu Val Phe Gln Pro Pro Ala

Asn

675 675 680

680 685 685

Gly Tyr Trp Val Gln Ala Pro Lys Gly Tyr Trp Val Gln Ala Pro Lys Thr Phe Thr Ile

Thr Phe Thr IIe Asn Phe Glu Asn Phe Glu Glu

Glu 690 695

690 695 700

700 Leu Glu Glu Ala Asn Gln Val Asn Gly Leu Tyr

Leu Glu Glu Ala Asn Gln Val His Tyr Glu Val Lys

Asn Gly Leu Tyr His Tyr Glu Val

Lys

705 710 705 710

715 720 715 720

Ile Asn Val Arg Asp Ile Ala Asn Ile Asn Val Arg Asp Ile Ala Asn Ile Gln Asp Asp

Ile Gln Asp Asp Thr Val Leu Arg Thr Val Leu Arg

725 725 730

730 735 735

Asn Met IIe Leu IIe Phe Ala Asp Asn Met IIe Leu IIe Phe Ala Asp Val Gln Ser Asp

Val Gin Ser Asp Phe Ala Gly Phe Ala Gly Arg

Arg

740 740 745

745 750 750

Val Phe Val Asp Asn Val Arg Val Phe Val Asp Asn Val Arg Phe Glu Ala Ser

Phe Glu Ala Ser Ala Thr Glu Ala Thr Glu Pro Val

Pro Val 755 760



760 765 Glu Pro Val Glu Pro Val Asp Pro Ala Pro Val Glu

Glu Pro Val Glu Pro Val Asp Pro Pro Glu Pro Val

Ala Pro Val Glu Pro Glu Pro Val

770 775 770 775

780 780

Asp Pro Gly Glu Glu Thr Pro Asp Pro Gly Glu Glu Thr Pro Pro Val Asp Glu

Pro Val Asp Glu Lys Glu Ala Ala Lys Glu Ala Ala Lys

Lys 785 790

785 790 795 800

795 800 Glu Glu Arg Glu Ala Ala Lys Ala Glu Arg Glu Ala

Glu Glu Arg Glu Ala Ala Lys Ala Ala Arg Glu Ala

Glu Arg Glu Ala Ala Arg Glu Ala

805 805 810

810 815 815

Ala Lys Glu Glu Arg Glu Glu Ala Ala Lys Glu Glu Arg Glu Glu Ala Arg Glu Ala Ala

Arg Glu Ala Ala Lys Glu Glu Arg Lys Glu Glu Arg

820 820 825

825 830 830

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Ala Ala Arg Glu Ala Ala Lys Ala Ala Ala Lys Ala

835 835 840

840 845 845

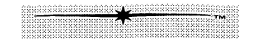
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Lys Lys Lys 850 855

850 855

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ile Prie Giu Giy(Trp)Sei Giri Lys Giy(Trp)Sei Giri Lys Vai

Val 5 10 15

10 15

Leu Ala Ala Glu Gly Asn Thr Arg Leu Ala Ala Glu Gly Asn Thr Arg Glu Asp Asn

Glu Asp Asn Phe Lys His Leu Phe Lys His Leu Leu

Leu 20 25

20 30

25 Gly Asn Asp Asn Val Lys Arg Pro Ser Glu Ala

Gly Asn Asp Asn Val Lys Arg Gly Ala Leu Gln Leu

Pro Ser Glu Ala Gly Ala Leu Gln 35 40

Leu 45

35

40 45

Gin Glu Val Asp Gly Gln Met Gin Glu Val Asp Gly Gln Met Thr Leu Val Asp

Thr Leu Val Asp Gln His Gly Glu Gln His Gly Glu Lys

Lys 50 55

50 55 60

60 Ile Gln Leu Arg Gly Met Ser Thr His Gly Leu Gln

Ile Gln Leu Arg Gly Met Ser Thr Trp Phe Pro Glu

His Gly Leu Gln Trp Phe Pro 65 70

Glu 75 80

__

65 70

75 80

Ile Leu Asn Asp Asn Ala Tyr Lys Ile Leu Asn Asp Asn Ala Tyr Lys Ala Leu Ser

Ala Leu Ser Asn Asp Trp Asp Asn Asp Trp Asp Ser

Ser 85 90

85 95

90 95 Asn Met Ile Arg Leu Ala Met Tyr Val Gly Glu Asn

Asn Met Ile Arg Leu Ala Met Tyr Gly His Ala Thr

Val Gly Glu Asn Gly His Ala Thr 100 105



Asn Pro Glu Leu IIe Lys Gln Arg Asn Pro Glu Leu IIe Lys Gln Arg Val IIe Asp Gly

Val Ile Asp Gly Ile Glu Leu Ala lie Glu Leu Ala

Ile Glu Asn Asp Met Tyr Val Ile Ile Glu Asn Asp Met Tyr Val Ile Val Asp Trp His

Val Asp Trp His Val His Ala Pro Val His Ala Pro

135 130

Gly Asp Pro Arg Asp Pro Val Tyr Gly Asp Pro Arg Asp Pro Val Tyr Ala Gly Ala Lys

Ala Gly Ala Lys Asp Phe Phe Asp Phe Phe Arg

Arg

150 155

Glu lle Ala Ala Leu Tyr Pro Asn Asn Pro His lle

Glu lle Ala Ala Leu Tyr Pro Asn Ile Tyr Glu Leu

Asn Pro His Ile Ile Tyr Glu Leu

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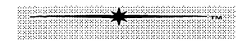
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Lys Glu Tyr Ala Asp Pro Ile Val

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Val

215 220



220 Gly Ser Pro Asn Trp Ser Gln Arg Pro Asp Leu

Gly Ser Pro Asn Trp Ser Gln Ala Ala Asp Asn Pro

230 Arg Pro Asp Leu Ala Ala Asp 225

Asn Pro

240 235

225 230

235 240

lle Asp Asp His His Thr Met Tyr Ile Asp Asp His His Thr Met Tyr Thr Val His Phe

Thr Val His Phe Tyr Thr Gly Ser Tyr Thr Gly Ser

250 245 245

255 255 250

His Ala Ala Ser Thr Glu Ser Tyr His Ala Ala Ser Thr Glu Ser Tyr Pro Pro Glu Thr

Pro Pro Glu Thr Pro Asn Ser Pro Asn Ser Glu

Glu 260 265

> 260 270

265 270

Arg Gly Asn Val Met Ser Asn Arg Gly Asn Val Met Ser Asn Thr Arg Tyr Ala

Thr Arg Tyr Ala Leu Glu Asn Gly Leu Glu Asn Gly Val

280 Val 275

285 275

280 285 Ala Val Phe Ala Thr Glu Trp Gly Thr Ser Gln Ala

Ala Val Phe Ala Thr Glu Trp Gly Asn Gly Asp Gly

295 Thr Ser Gln Ala Asn Gly Asp 290

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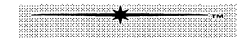
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Asn Asn Ile Ser Trp Ala Asn Trp Asn Lys Asn Glu Vai

330 Ser Leu Thr Asn Lys Asn Glu 325

335 Val



330 335

Ser Gly Ala Phe Thr Pro Phe Ser Gly Ala Phe Thr Pro Phe Glu Leu Gly Lys

Glu Leu Gly Lys Ser Asn Ala Ser Asn Ala Thr Ser

Thr Ser 340 345

340 350

345 350 Leu Asp Pro Gly Pro Asp Gln Val Trp Val Pro

Leu Asp Pro Gly Pro Asp Gln Glu Glu Leu Ser Leu

Val Trp Val Pro Glu Glu Leu Ser 355 360

Leu 365

355

360 365

Ser Gly Glu Tyr Val Arg Ala Arg Ser Gly Glu Tyr Val Arg Ala Arg Ile Lys Gly Val

Ile Lys Gly Val Asn Tyr Glu Pro Asn Tyr Glu Pro

370 375 370 375

380 380

Ile Asp Arg Thr Lys Tyr Thr Lys Ile Asp Arg Thr Lys Tyr Thr Lys Val Leu Trp Asp

Val Leu Trp Asp Phe Asn Asp Phe Asn Asp Gly

Gly 385 390

385 390 395 400

395 400

Thr Lys Gln Gly Phe Gly Val Thr Lys Gln Gly Phe Gly Val Asn Ser Asp Ser

Asn Ser Asp Ser Pro Asn Lys Pro Asn Lys Glu Leu

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Ile Ala Val Asp Asn Glu Asn Asn Ser Gly Leu Asp

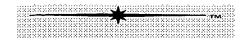
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Asp 430

420

425 430

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Gly Asn Phe Trp Ala Asn Ala Ala Asn Ala Arg Leu

Arg Leu 435 440

435 445

440 445 Ser Ala Asn Gly Trp Gly Lys Ser Val Asp Ile Leu

Ser Ala Asn Gly Trp Gly Lys Ser Gly Ala Glu Lys

Val Asp IIe Leu Gly Ala Glu Lys 450 455

450 455 460

460

Leu Thr Met Asp Val Ile Val Asp Leu Thr Met Asp Val Ile Val Asp Glu Pro Thr Thr

Glu Pro Thr Thr Val Ala Ile Ala Val Ala Ile Ala

465 470 465 470

475 480 475 480

Ala Ile Pro Gln Ser Ser Lys Ser Ala Ile Pro Gln Ser Ser Lys Ser Gly Trp Ala Asn

Gly Trp Ala Asn Pro Glu Arg Ala Pro Glu Arg Ala

485 485 490

490 495 495

Val Arg Val Asn Ala Glu Asp Val Arg Val Asn Ala Glu Asp Phe Val Gln Gln

Phe Val Gln Gln Thr Asp Gly Thr Asp Gly Lys Tyr

Lys Tyr 500 505

500 510

505 Lys Ala Gly Leu Thr Ile Thr Gly Glu Asp Ala Pro

Lys Ala Gly Leu Thr Ile Thr Gly Ser Leu Glu Ala

Glu Asp Ala Pro Ser Leu Glu 515 520

Ala 525

515

520 525

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Thr lie Asn Asn lie lie Leu Phe lie lie Leu Phe

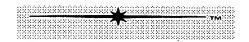
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540 540

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lle Tyr Leu Asp Thr lle Lys Val Thr lle Lys Val

545 550 545 550



555 560 555 560

lle Gly Pro Glu Val Glu lle Pro lle Gly Pro Glu Val Glu lle Pro Val Val His Asp

Val Val His Asp Pro Lys Gly Glu Pro Lys Gly Glu

565 565 570

570 575

Ala Val Leu Pro Ser Val Phe Glu Ala Val Leu Pro Ser Val Phe Glu Asp Gly Thr

Asp Gly Thr Arg Gln Gly Trp Arg Gln Gly Trp Asp

Asp 580 585

580 590

585 590

Trp Ala Gly Glu Ser Gly Val Lys Trp Ala Gly Glu Ser Gly Val Lys Thr Ala Leu Thr

Thr Ala Leu Thr Ile Glu Glu Ala Ile Glu Glu Ala

595 595 600

600 605 605

Asn Gly Ser Asn Ala Leu Ser Asn Gly Ser Asn Ala Leu Ser Trp Glu Phe Gly

Trp Glu Phe Gly Tyr Pro Glu Val Tyr Pro Glu Val Lys

Lys 610 615

610 615 620

620

Pro Ser Asp Asn Trp Ala Thr Ala Pro Ser Asp Asn Trp Ala Thr Ala Pro Arg Leu

Pro Arg Leu Asp Phe Trp Lys Asp Phe Trp Lys Ser

Ser 625 630

625 630 635 640

635 640 Asp Leu Val Arg Gly Glu Asn Asp Tyr Val Thr

Asp Leu Val Arg Gly Glu Asn Phe Asp Phe Tyr Leu

Asp Tyr Val Thr Phe Asp Phe 645 650

Tyr Leu 655

645

650 655

Asp Pro Val Arg Ala Thr Glu Gly Asp Pro Val Arg Ala Thr Glu Gly Ala Met Asn Ile

Ala Met Asn Ile Asn Leu Val Asn Leu Val Phe

Phe 660 665



665 Gln Pro Pro Thr Asn Gly Tyr Trp Val Gln Ala Pro

Gln Pro Pro Thr Asn Gly Tyr Trp Lys Thr Tyr Thr

Val Gln Ala Pro Lys Thr Tyr Thr 675 680

675 685

680 685

Ile Asn Phe Asp Glu Leu Glu Ile Asn Phe Asp Glu Leu Glu Glu Ala Asn Gln

Glu Ala Asn Gln Val Asn Gly Val Asn Gly Leu Tyr

Leu Tyr 690 695

690 695 700

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715 720

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Ser 725 730

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Thr Pro Pro Val Asp Glu Lys Glu Ala

Glu Ala Lys Lys Glu Gln Lys Glu 770 775

Ala 780

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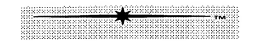
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Met

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Leu

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621

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His Leu Leu Gly Asn Glu Asn 35 Val Lys

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110 105

Asn Ala



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140 145 140 145

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Val Ile Val Asp Trp His Val His Pro Arg Asp Pro

Ala Pro Gly Asp Pro Arg Asp

Pro

150 150 155

155 160 160

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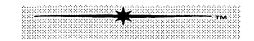
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Ser 190

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aaa 1101

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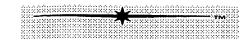
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Asp Asn Pro Ile Asn Asp His 245 250

His Thr 255



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290 295

300 305

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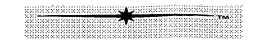
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Asn Glu Val Ser Gly Ala Phe

Thr Pro

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Ala Thr Ser Leu Asp Pro Gly 355 360

Pro Asp 365

355 360

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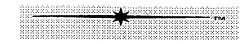
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1677



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Lys Glu Ala Ile Glu Val Glu Asn 420 425

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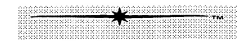
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Val Asp Glu Pro Thr Thr Val Ala Pro Gln Ser Thr

Ile Ala Ala Ile Pro Gin Ser Thr 485 490

485 495

490 495

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Lys His Gly Trp Ala Asn Pro Glu Val Thr Glu Ala

Arg Ser Val Lys Val Thr Glu Ala 500 505

500 510

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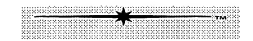
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550 560

555 560

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Lys Val Thr Gly Lys Ile Val Glu 565 570

565 575

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2253

Ile Pro Val Val His Ser Pro Lys Ile Pro Val Val His Ser Pro Lys Gly Asp Ala Ala Gly Asp Ala Ala Leu Pro Ser Leu Pro Ser Asn

Asn 580 585

580 590

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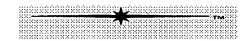
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Phe Tyr Ile Asp Pro Ala Arg Ala Thr

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Gly 685

675 680

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Phe Thr Ile Asn Phe Glu Glu 690 695

Leu 700 705

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ggg tta tac cat tat gaa gtg aaa

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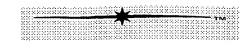
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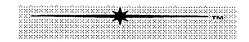
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795 800 800

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820 830

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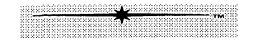
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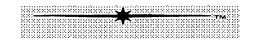
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Gly Asn

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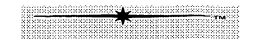
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Gly Leu Gln Trp Phe Pro Glu lle 70 75

Leu 80

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Asn Met 85 90

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Pro

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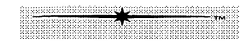
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Trp His Val His Ala Pro Gly Asp 135

135 145

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Ile 150 155

150 160

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gca gct tta tac cct aat aat cca tta gcg aat 826

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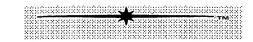
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Asn Asn 320

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315 320

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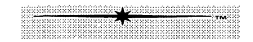
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340 345 Leu Asp

340 345 350

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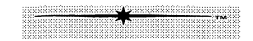
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445 450

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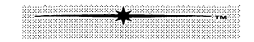
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490 495 495

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Gln Gln Thr Asp Gly Lys Tyr Lys 500 505

Ala 510

500 505

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Ala Pro Ser Leu Glu Ala Ile Ala 515 520

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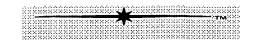
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540 545 545



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1978

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His Asp Pro Lys Gly Glu Ala Val 565

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2074

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Gly Thr Arg Gln Gly Trp Asp Trp 580

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Ala

590

580

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590

gaa gca aac ggt

2122

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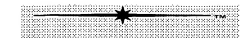
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2170

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Phe Gly Tyr Pro Glu Val Lys Pro Ser

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620 625 625

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Leu Asp Phe Trp Lys Ser Asp 630 635

Leu 640

630

635 640

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Thr Phe Asp Phe Tyr Leu Asp 645 650

Pro 655

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650 655 gta ttc cag cca 2314

gtt cgt gca aca gaa ggc gca atg aat atc aat tta gta ttc cag cca

2314

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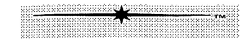
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2362 Tyr Thr Ile Asn

Pro Thr Asn Gly Tyr Trp Val Gln Ala Pro Lys Thr Tyr Thr Ile Asn



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Phe Asp Glu Leu Glu Glu Ala Gly Leu Tyr His Tyr

Asn Gln Val Asn Gly Leu Tyr 695 700

His Tyr 705

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700 705

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Ile Thr Asn Ile Gln Asp Asp Thr 710 715

710 720

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tta cta cgt aac atg atg atc att ttt agt gac ttt 2506

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Phe 725 730

725 735

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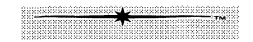
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Thr

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Pro Val Asp Glu Lys Glu Ala Lys Glu Ala Glu Lys

Lys Glu Gln Lys Glu Ala Glu Lys 775 780

775 785

780 Gaa gag aaa gaa gaa gaa gaa gaa aag aaa

gaa gag aaa gaa gca gta aaa gaa gct aaa gaa gaa 2698

gaa gaa aag aaa gaa gct aaa

gaa gaa 2698

Glu Glu Lys Glu Ala Val Lys Glu Glu Glu Lys Glu Ala Val Lys Glu Glu Lys Lys Glu

Glu Lys Lys Glu Ala Lys Glu Glu Ala Lys Glu Glu

790 790 795

795 800 800

aag aaa gca atc aaa aat gag Aag aaa gca atc aaa aat gag gct acg aaa aaa

gct acg aaa aaa taatctatta taatctatta aactagttat 2751

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Lys Lys Ala lle Lys Asn Glu Ala

Thr Lys Lys

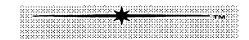
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2783



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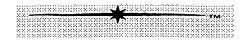
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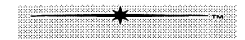
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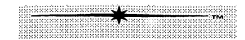
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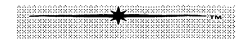
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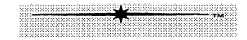
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based on S131b cellulase and cellulase and KSM-64 cellulase



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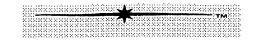
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ggcttgtgct ggtcgaccca actgc Ggcttgtgct ggtcgaccca actgc 25 25

【図面の簡単な説明】

[BRIEF DESCRIPTION OF THE DRAWINGS]

【図1】

Hの影響を示す図である。

[FIG. 1]

本発明のアルカリセルラーゼ It is the figure showing the influence of pH (N 1 3 1 a) 活性に及ぼす p which affects the alkali cellulase (N131a) activity of this invention.

【図2】

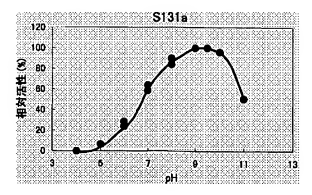
Hの影響を示す図である。

[FIG. 2]

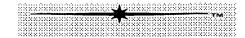
本発明のアルカリセルラーゼ It is the figure showing the influence of pH (N 1 3 1 b) 活性に及ぼす p which affects the alkali cellulase (N131b) activity of this invention.

【図1】

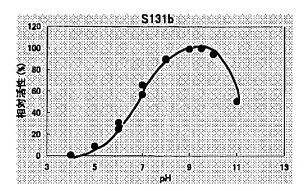
[FIG. 1]



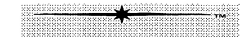
相対活性: Relative activity



[図2]



相対活性: Relative activity



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